

Office de la Propriété Intellectuelle du Canada

Un organisme d'Industrie Canada Canadian Intellectual Property Office

An agency of Industry Canada CA 2433626 A1 2002/07/18

(21) 2 433 626

(12) DEMANDE DE BREVET CANADIEN CANADIAN PATENT APPLICATION

(13) A 1

対な特別2003-294397

(86) Date de dépôt PCT/PCT Filing Date: 2001/12/27

(87) Date publication PCT/PCT Publication Date: 2002/07/18

(85) Entrée phase nationale/National Entry: 2003/06/27

(86) N° demande PCT/PCT Application No.: JP 2001/011569

(87) N° publication PCT/PCT Publication No.: 2002/055751

(30) Priorités/Priorities: 2000/12/29 (2000-404991) JP; 2001/03/30 (2001-102186) JP

(51) Cl.Int.7/Int.Cl.7 C22C 38/00, C21D 9/46

(71) Demandeur/Applicant: NIPPON STEEL CORPORATION, JP

(72) Inventeurs/Inventors: TAKADA, YOSHIHISA, JP; SUEHIRO, MASAYOSHI, JP; SENUMA, TAKEHIDE, JP

(74) Agent: FETHERSTONHAUGH & CO.

(54) Titre: PLAQUE D'ACIER A PLACAGE EN ZINC MOULE A HAUTE RESISTANCE POSSEDANT UNE EXCELLENTE ADHESION EN DEPOT ET PARFAITEMENT ADAPTEE AU FORMAGE A LA PRESSE ET PROCEDE DE FABRICATION ASSOCIE

(54) Title: HIGH-STRENGTH MOLTEN-ZINC-PLATED STEEL PLATE EXCELLENT IN DEPOSIT ADHESION AND SUITABILITY FOR PRESS FORMING AND PROCESS FOR PRODUCING THE SAME

(57) Abrégé/Abstract:

A high-strength molten-zinc-plated steel plate excellent in suitability for press forming and in deposit adhesion which is useful as automotive, building, electrical, and other members. The plated steel plate comprises: (a) a steel plate base which contains 0.05 to 0.2 wt.% carbon, 0.2 to 2.0 wt.% silicon, 0.2 to 2.5 wt.% manganese, and 0.01 to 1.5 wt.% aluminum, the silicon and aluminum satisfying the relationship 0.4 (%) \leq Si+0.8Al (%) \leq 2.0%, and contains at least one member selected from the group consisting of (i) 0.003 to 1.0 wt.% tin, (ii) at least one of antimony, bismuth, and selenium in a total amount of 0.005 to 1.0 wt.%, (iii) at least one of beryllium, magnesium, calcium, and zirconium in a total amount of 0.005 to 1.0 wt.%, and (iv) at least one of scandium, yttrium, lanthanum, and cerium in a total amount of 0.005 to 1.0 wt.%, and in which the remainder consists of iron and unavoidable impurities and the content of residual austenites in the steel structure is 2 to 20 vol.%; and (b) a zinc deposit layer formed on the steel plate base.





ABSTRACT

Disclosed are a high strength hot-dip galvanized or galvannealed steel sheet, which has improved press formability and plating adhesion and is useful as a member for automobile, building, electric or other members, and a process for producing the same.

5

10

15

20

25

This high strength hot-dip galvanized or galvannealed steel sheet comprises: (a) a steel sheet substrate comprising, by weight, carbon (C): 0.05 to 0.2%, silicon(Si):0.2 to 2.0%, manganese (Mn): 0.2 to 2.5%, and aluminum (Al): 0.01 to 1.5%, the silicon and the aluminum having a mutual relationship represented by formula 0.4 (%) ≤ Si + 0.8Al (%) ≤ 2.0 (%), the steel sheet substrate further comprising at least one member selected from the group consisting of (i) 0.003 to 1.0% of tin (Sn), (ii) 0.005 to 1.0% in total of at least one member selected from antimony (Sb), bismuth (Bi), and selenium (Se), (iii) 0.005 to 1.0% in total of at least one member selected from beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr), and (iv) 0.005 to 1.0% in total of at least one member selected from scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce), with the balance consisting of iron (Fe) and unavoidable impurities, the volume fraction of retained austenite in the steel structure being 2 to 20%; and (b) a zinc (Zn) coating layer provided on said steel sheet substrate.

HIGH STRENGTH HOT-DIP GALVANIZED OR GALVANNEALED STEEL SHEET
HAVING IMPROVED PLATING ADHESION AND PRESS FORMABILITY AND
PROCESS FOR PRODUCING THE SAME

[BACKGROUND OF THE INVENTION]

Field of the Invention

5

10

15

20

25

30

35

The present invention relates to a high strength steel sheet useful as automobile, building, electric or other members and a process for producing the same. More particularly, the present invention relates to a high strength hot-dip galvanized steel sheet which has improved bulging formability at the time of press forming and improved plating adhesion and a process for producing the same. The term "high strength hot-dip galvanized or galvannealed steel sheet" as used herein includes high strength hot-dip galvanized steel sheets (GI) and high strength hot-dip galvannealed steel sheets (GA).

Background Art

To meet a demand for lower fuel consumption of automobiles and the like in recent years, studies have been made on a weight reduction of members such as cross members and side members of automobiles and the like. Regarding materials, studies on an increase in strength have been made from the viewpoint of ensuring strength even in reduced thickness of members.

In general, however, increasing the strength of materials results in deteriorated press formability of the materials. Therefore, in order to realize the weight reduction of the above members, steel sheets capable of simultaneously satisfying good press formability and high strength are required, and the development of such steel sheets has been desired in the art.

Among index values for formability are included elongation and, in addition, n value and r value in tensile tests. A task to be accomplished in recent years is to simplify the step of pressing through one-piece molding. Therefore,

what is particularly important is that the n value corresponding to uniform elongation is large.

To this end, hot rolled steel sheets or cold rolled steel sheets, which utilize transformation induced plasticity of retained austenite present in steel, have been developed. In these steel sheets, only about 0.07 to 0.4% of carbon (C), about 0.3 to 2.0% of silicon (Si), and about 0.2 to 2.5% of manganese (Mn) are used as basic alloying elements without incorporating any expensive alloying element, and heat treatment characterized by, after annealing in a two-phase region, performing bainite transformation at a temperature of about 300 to 450°C is carried out to allow retained austenite to exist in the metallurgical structure. These steel sheets are disclosed, for example, in Japanese Patent Laid-Open Nos. 230715/1989 and 217425/1990.

Steel sheets, to which the above technique can be applied, are not limited to cold rolled steel sheets produced by continuous annealing. For example, Japanese Patent Laid-Open No. 79345/1989 discloses that this technique can also be applied to hot rolled steel sheets by controlling cooling temperature of a coiling or a runout table.

The production of automobiles of higher grade has led to a tendency toward coating of automobile members with a view to improving corrosion resistance and appearance. At the present time, galvanized steel sheets are used in most of the automobile members except for specific members mounted within the automobiles. Accordingly, from the viewpoint of corrosion resistance, coating which is useful for these steel sheets is hot-dip galvanizing, or hot-dip galvannealing wherein, after hot-dip galvanizing, alloying treatment is carried out. Among these high-tensile steel sheets, steel sheets having a high silicon content are likely to form an oxide film on the surface thereof. This poses problems including that very small non-coated sites occur in hot-dip galvanizing, or plating adhesion of the worked part after the completion of alloying is deteriorated. Thus, at the

present time, any high-silicon-base high-tensile and high-ductile hot-dip galvannealed steel sheets possessing excellent plating adhesion of the worked part and, at the same time, excellent corrosion resistance have not been put into practical use.

5

10

15

20

25

30

35

In the steel sheets disclosed, for example, in Japanese Patent Laid-Open Nos. 230715/1989 and 217425/1990, however, it should be noted that 0.3 to 2.0% of silicon is contained and the retained austenite is ensured by utilizing unique bainite transformation of the steel. Therefore, the contemplated metallurgical structure cannot be provided without considerably close control of cooling after annealing in an intercritical temperature range or holding in a temperature range of about 300 to 450°C, and strength and elongation disadvantageously deviate from the target range.

This heat history can be realized on a commercial scale by continuous annealing equipment or a runout table after hot rolling and the step of coiling. Since, however, the transformation of austenite is rapidly completed at 450 to 600°C, control should be carried out so that the residence time is short particularly at 450 to 600°C. Further, even at 350 to 450°C, the metallurgical structure undergoes a significant change depending upon the holding time. Therefore, when the heat treatment conditions have deviated from predetermined conditions, only unsatisfactory strength and elongation are provided.

Further, since the residence time at 450 to 600°C is long and a large amount of silicon, which deteriorates suitability for coating, is contained as an alloying element, the production of a coated steel sheet by passing the steel sheet through hot-dip galvanizing equipment is impossible. After all, poor surface corrosion resistance of the steel sheet containing about 0.3 to 2.0% of silicon disadvantageously hinders the widespread commercial use of the steel sheets.

In order to solve the above problem, for example,

Japanese Patent Laid-Open Nos. 247586/1993 and 145788/1994 disclose steel sheets having suitability for coating improved by regulating the content of silicon. In the technique disclosed in the above publications, retained austenite is formed by adding aluminum (Al) instead of silicon. However, as with silicon, aluminum is more likely to be oxidized than iron (Fe). Specifically, aluminum is likely to form an oxide film on the surface of the steel sheet. Therefore, disadvantageously, satisfactory plating adhesion cannot be ensured.

Further, for example, Japanese Patent Laid-Open Nos. 333552/1992 and 346644/1992 disclose a hot-dip galvannealing method for a high-silicon-base high-strength steel sheet. In this method, after pre-coating of nickel (Ni), the pre-coated steel sheet is rapidly heated at a low temperature and is subjected to hot-dip galvanizing, followed by alloying treatment. Since, however, this method requires nickel pre-coating, disadvantageously, equipment for nickel pre-coating should be newly provided.

20

25

30

35

5

10

15

[SUMMARY OF THE INVENTION]

The present inventors have made studies on the solution of the above problems of the prior art and on an improvement in surface corrosion resistance and, as a result, have found the chemical composition and characteristics of the metallurgical structure of a high strength steel sheet which can be produced even in hot-dip galvanizing equipment and has good press formability.

An object of the present invention is to solve the above problems of the prior art and to provide a high strength hot-dip galvanized or galvannealed steel sheet possessing good press formability and plating adhesion and a production process which can efficiently produce this steel sheet.

The present inventors have made extensive and intensive studies on the relationship between suitability for coating and steel constituents with a view to providing a high strength

hot-dip galvanized or galvannealed steel sheet and a production process thereof which can attain the above object of the present invention. This has led to the completion of the present invention.

- (1) According to the present invention, there is provided a high strength hot-dip galvanized or galvannealed steel sheet having improved plating adhesion and press formability, said steel sheet comprising:
 - (a) a steel sheet substrate comprising, by weight, carbon (C): 0.05 to 0.2%,

silicon (Si): 0.2 to 2.0%,

10

manganese (Mn): 0.2 to 2.5%, and

aluminum (Al): 0.01 to 1.5%,

said silicon and said aluminum having a mutual 15 relationship represented by formula

 $0.4 (%) \le Si + 0.8Al (%) \le 2.0%,$

said steel sheet substrate further comprising at least one member selected from the group consisting of

- (i) 0.003 to 1.0% of tin (Sn),
- 20 (ii) 0.005 to 1.0% in total of at least one member selected from antimony (Sb), bismuth (Bi), and selenium (Se),
 - (iii) 0.005 to 1.0% in total of at least one member selected from beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr), and
- (iv) 0.005 to 1.0% in total of at least one member selected from scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce), with the balance consisting of iron (Fe) and unavoidable impurities, the volume fraction of retained austenite in the steel structure being 2 to 20%; and
- 30 (b) a zinc (Zn) coated layer provided on said steel sheet substrate.
- (2) In a preferred embodiment of the present invention, said steel sheet substrate further comprises, by weight, at least one member selected from nickel (Ni): not more than 2.0%, copper (Cu): not more than 2.0%, and cobalt (Co): less than 0.3%.

(3) In a preferred embodiment of the present invention, said steel sheet substrate further comprises, by weight, at least one member selected from molybdenum (Mo): less than 0.5%, chromium (Cr): less than 1.0%, vanadium (V): less than 0.3%, titanium (Ti): less than 0.06%, niobium (Nb): less than 0.06%, and boron (B): less than 0.01%.

5

10

15

20

25

30

35

- (4) In a preferred embodiment of the present invention, said zinc coated layer is a galvannealed layer comprising zinc (Zn): 80 to 91%, iron (Fe): 8 to 15%, and aluminum (Al): not more than 1%.
- (5) In a preferred embodiment of the present invention, said zinc coated layer is a galvanized layer comprising zinc: not less than 80% and aluminum: not more than 1%.
- (6) According to the present invention, there is provided a process for producing the above hot-dip galvanized or galvannealed steel sheet having a galvannealed layer of the present invention, said process comprising the steps of:

providing a cold rolled steel sheet having the above chemical composition of the steel sheet substrate;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, and subjecting the cooled steel sheet to hot-dip galvannealing process; and then

holding the coated steel sheet in the temperature range of 450 to 600°C for 5 sec to 2 min and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

(7) According to the present invention, there is provided another process for producing the above hot-dip galvanized or galvannealed steel sheet having a hot-dip galvannealed layer of the present invention, said process comprising the steps of:

providing a cold rolled steel sheet having the above chemical composition of the steel sheet substrate;

annealing the cold rolled steel sheet for 10 sec to 6

min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, holding the cooled steel sheet in said cooling temperature range for not more than 10 min, and then subjecting the cooled steel sheet to hot-dip galvannealed process; and then

5

10

15·

20

25

30

35

holding the coated steel sheet in the temperature range of 450 to 600°C for 5 sec to 2 min and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

(8) According to the present invention, there is provided a process for producing the above hot-dip'galvanized or galvannealed steel sheet having a galvanized layer of the present invention, said process comprising the steps of:

providing a cold rolled steel sheet having the above chemical composition of the steel sheet substrate;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, and subjecting the cooled steel sheet to hot-dip galvanizing process; and then

cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec .

(9) According to the present invention, there is provided another process for producing the above hot-dip galvanized or galvannealed steel sheet having a galvanized layer of the present invention, said process comprising the steps of:

providing a cold rolled steel sheet having the above chemical composition of the steel sheet substrate;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, holding the cooled steel sheet in said cooling temperature range for not more than 10 min, and then subjecting the cooled steel sheet to hot-dip galvanizing process; and then

cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

[DETAILED DESCRIPTION OF THE INVENTION]

5 (a) Steel sheet substrate

10

15

20

25

30

35

In the steel sheet substrate according to the present invention, the chemical composition is limited from the viewpoint of providing high strength hot-dip galvanized or galvannealed steel sheets possessing good press formability and good plating adhesion. The reasons for the limitation of the chemical composition will be described in detail.

Basic constituents

Carbon (C) is an austenite stabilizer. In the intercritical temperature range and in the bainite transformation temperature range, carbon migrates from ferrite and is enriched in austenite. As a result, even after cooling to room temperature, 2 to 20% of chemically stabilized austenite is present and improves formability through transformation induced plasticity. If the content of carbon is less than 0.05%, then it is difficult to ensure not less than 2% of retained austenite, making it impossible to attain the contemplated effect. On the other hand, a carbon content exceeding 0.2% should be avoided, because the weldability is deteriorated.

Silicon (Si) does not dissolve in cementite and suppresses the precipitation of cementite. By virtue of this action, the transformation from austenite to cementite at 350 to 600°C is delayed, during which time the enrichment of carboninaustenite is promoted. This can enhance the chemical stability of austenite, causes transformation induced plasticity, and thus enables retained austenite, which can improve the formability, to be ensured. If the content of silicon is less than 0.2%, then the effect of silicon cannot be attained. On the other hand, when the silicon content is excessively high, the suitability for coating is deteriorated. Therefore, the silicon content should be not more than 2.0%.

5

10

15

20

25

30

35

Manganese (Mn) is an austenite former. manganese can prevent the decomposition of austenite to pearlite in the course of cooling to 350 to 600°C after annealing in the intercritical temperature range. Therefore, manganese acts so that retained austenite is contained in the metallurgical structure after cooling to room temperature. When the content of manganese is less than 0.2%, in order to prevent the decomposition of austenite to pearlite, the cooling rate should be increased to such a level that could not be controlled on a commercial scale. On the other hand, when the manganese disadvantageous. content exceeds 2.5%, the formation of a banded structure is significant, resulting in deteriorated properties of the steel sheet. Further, in this case, upon spot welding, the spot weld zone is disadvantageously likely to be broken within the nugget. Furthermore, increasing the manganese content deteriorates the suitability for coating.

Aluminum (Al) is used as a deoxidizer. At the same time, as with silicon, aluminum does not dissolve in cementite and, in holding the steel sheet at 350 to 600°C, suppresses the precipitation of cementite and delays the transformation from austenite to cementite. Since, however, aluminum has higher ferrite forming ability than silicon, the transformation starts in an earlier stage. In this case, even when the holding time is very short, carbon is enriched in austenite from the start of annealing in the intercritical temperature range. This can enhance the chemical stability, and thus, the amount of martensite, which deteriorates the formability, present in the metallurgical structure after cooling to room temperature is very small. Therefore, the coexistence of aluminum and silicon can reduce a change in strength and elongation depending upon conditions for holding at 350 to 600°C, and steel sheets having a combination of high strength with good press formability can be easily provided. attaining the above effect, the addition of aluminum in an amount of not less than 0.01% is necessary. The addition of

aluminum in an amount of not less than 0.1% is preferred. In addition, aluminum, together with silicon, should be added so that the content of "Si + 0.8Al" is not less than 0.4%. On the other hand, as with the case of silicon, an aluminum content exceeding 1.5% deteriorates plating adhesion and thus should be avoided. Furthermore, from the viewpoint of ensuring the plating adhesion, aluminum, together with silicon, should be added so that the content of "Si + 0.8Al" is not more than 2.0%.

Selective basic constituents

5

10

15

20

25

30

35

Tin (Sn), antimony (Sb), bismuth (Bi), selenium (Se), beryllium (Be), magnesium (Mg), calcium (Ca), zirconium (Zr), scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) are elements which are most important to the present invention. The addition of at least one of these elements can improve the wettability and the plating adhesion of the steel sheet by hot-dip galvanizing or galvannealing. As a result, steel sheets having excellent suitability for coating and formability can be produced.

In steel sheets containing silicon and aluminum, when coated steel sheets are produced in a continuous hot-dip galvanizing line, oxides of silicon and aluminum are formed on the surface of the steel sheets, resulting in deteriorated plating adhesion. However, the addition of at least one of the above elements can improve the suitability of the steel sheets for coating.

In a preferred embodiment of the present invention, tin is added in an amount of 0.003 to 1.0%. In steel sheets containing silicon and aluminum, when coated steel sheets are produced in a continuous hot-dip galvanizing line, oxides of silicon and aluminum are formed on the surface of the steel sheets and deteriorate the plating adhesion to the steel sheets. However, since tin is an element which is less likely to be oxidized than iron and, at the same time, is likely to segregate on the surface of steel sheets, tin is enriched in the surface layer of the steel sheets to suppress the formation of oxides

of silicon and aluminum, whereby the deterioration in plating adhesion is prevented. When the content of tin is less than 0.003%, satisfactory plating adhesion cannot be provided in the steel of the present invention. When more significant development of the above effect is desired, the addition of tin in an amount of not less than 0.005% is preferred. The amount of tin added is more preferably not less than 0.008%. On the other hand, when tin is added in an amount of more than 1.0%, cracking disadvantageously occurs at the time of hot rolling. As a result, good appearance of coating cannot be ensured. When better appearance of coating is desired, the addition of tin in an amount of not more than 0.5% is preferred.

5

10

In another preferred embodiment of the present invention, 15 0.005 to 1.0% in total of at least one of antimony, bismuth, and selenium is added. Antimony, bismuth, and selenium are likely to cause surface segregation and thus are enriched in the surface layer of the steel sheet to suppress the formation of oxides of silicon and aluminum. Consequently, even in the 20 case of high-silicon and/or high-aluminum steel, deterioration in plating adhesion can be prevented. effect can be attained by adding at least one of antimony, bismuth, and selenium. When the total amount of antimony, bismuth, and selenium is not less than 0.005%, satisfactory 25 plating adhesion can be provided. When a further improved effectis desired, the addition of at least two of these elements in a total amount of not less than 0.008% is preferred. addition of more than 1.0% in total of at least one of antimony, bismuth, and selenium causes surface segregation of these 30 elements in an excessive amount. As a result, good appearance of coating cannot be ensured. In order to maintain good appearance of coating, the addition of not more than 0.5% in total of at least one of antimony, bismuth, and selenium is preferred.

As with antimony, bismuth, and selenium, arsenic (As), tellurium (Te), polonium (Po), and germanium (Ge) can improve

5

10

15

20

25

30

35

suitability of the steel sheet for coating. Since, however, arsenic (As), tellurium (Te), polonium (Po), and germanium (Ge) are toxicelements and are very high in cost, these elements are excluded from the additive elements in the present invention.

In still another preferred embodiment of the present invention, 0.005 to 1.0% in total of at least one of beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr) is added. Since beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr) are very likely to form oxides, they can suppress the formation of silicon oxide and/or aluminum oxide which deteriorate suitability of high-silicon steel and/or high-aluminum steel for coating. This contributes to an improvement in suitability for coating. This effect can be attained by adding at least one of beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr). The addition of not less than 0.005% in total of at least one of beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr) can provide satisfactory plating adhesion. When a further improved effectisdesired, the addition of atleast two of these elements in a total amount of not less than 0.008% is preferred. addition of more than 1.0% in total of at least one of beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr) results in an increased amount of the formation of oxides of these elements and consequently makes it impossible to ensure good appearance of coating.

In a further preferred embodiment of the present invention, 0.005 to 1.0% in total of at least one of scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) is added. Since scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) are also likely to form oxides, they can suppress the formation of silicon oxide and/or aluminum oxide which deteriorate suitability of high-silicon steel and/or high-aluminum steel for coating. This contributes to an improvement in suitability for coating. Further, at the time of oxidation, scandium (Sc), yttrium (Y), lanthanum (La),

and cerium (Ce) vigorously act on surface irregularities of This contributes to improved plating the steel sheet. This effect can be attained by adding at least one adhesion. of scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce). The addition of not less than 0.005% in total of at least one of scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) can provide satisfactory plating adhesion. When a further improved effect is desired, the addition of at least two of these elements in a total amount of not less than 0.008% is preferred. The addition of more than 1.0% in total of at least one of scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) results in the formation of an increased amount of oxides of these elements and consequently makes it impossible to ensure good appearance of coating.

5

10

15

20

25

30

35

As with scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce), rare earth elements such as neodymium (Nd), gadolinium (Gd), and dysprosium (Dy) can improve the suitability for coating. These elements, however, are very high in cost and thus are excluded from additive elements in the present invention.

Further, the addition of a combination of two or more members, which are different from each other or one another in effect, selected from the group consisting of (i) tin (Sn), (ii) at least one member selected from antimony (Sb), bismuth (Bi), and selenium (Se), (iii) at least one member selected from beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr), and (iv) at least one member selected from scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce) can ensure better suitability for coating.

The addition of not less than 0.005% in total of at least one of these elements can provide satisfactory plating adhesion. When the total amount of at least one of these elements exceeds 1.0%, good appearance of coating cannot be ensured.

Optional constituents

In the steel sheet according to the present invention,

the above-described elements constitute basic constituents. In addition to these elements and iron, for example, at least one of nickel (Ni), copper (Cu), and cobalt (Co), which are austenite formers and, at the same time, can improve strength and plating adhesion, may be added. Also, at least one of molybdenum (Mo), chromium (Cr), vanadium (V), boron (B), titanium (Ti), niobium (Nb), and boron (B), which are hardenability improving elements (constituent (a) group), and/or at least one of REM (rare earth metal), calcium (Ca), zirconium (Zr), and magnesium (Mg), which reduce inclusions (constituent (b) group), may be added optionally together with at least one of the above-described elements, that is, nickel, copper, and cobalt, to the above basic constituents.

5

10

15

20

25

30

35

The reasons for the limitation of the contents of the above elements in the steel sheet will be described in detail.

Nickel (Ni), copper (Cu), and cobalt (Co), as with tin (Sn), are less likely to be oxidized than iron and thus are enriched on the surface of the steel sheet at the time of annealing to suppress the formation of oxides of silicon (Si), aluminum (Al) and the like which inhibit plating adhesion. Further, nickel (Ni), copper (Cu), and cobalt (Co), as with manganese (Mn), are austenite formers and, at the same time, as with silicon (Si) and aluminum (Al), do not dissolve in cementite. Therefore, in holding the steel sheet at 350 to 600°C, nickel (Ni), copper (Cu), and cobalt (Co) suppress the precipitation of cementite and delay the progress of transformation. Therefore, the addition of at least one of nickel (Ni), copper (Cu), and cobalt (Co) can provide better steel sheets. When nickel is added in an amount exceeding 2.0%, the contemplated effect is saturated. For this reason, the upper limit of the nickel content is 2.0%. When copper (Cu) is added in an amount exceeding 2.0%, the quality of the steel sheet is deteriorated as a result of precipitation of copper (Cu). For this reason, the upper limit of the copper content is 2.0%. Since cobalt (Co) is an expensive metal, the upper limit of the cobalt content is 0.3%. When tin and

5

10

15

20

25

30

35

copper (Co) are added in combination, preferably, tin and copper satisfy a mutual relationship represented by formula "Sn (%) + Cu (%) $< 3 \times Ni$ (%)" from the viewpoint of preventing hot cracking caused by tin and copper.

Molybdenum (Mo), chromium (Cr), vanadium (V), titanium (Ti), niobium (Ni), and boron (B) are strength improving elements, and REM, calcium (Ca), zirconium (Zr), and magnesium (Mg) are elements which combine with sulfur in the steel to reduce inclusions, thereby ensuring good elongation of the Preferably, the steel sheet substrate further comprises molybdenum (Mo): less than 0.5%, chromium (Cr): less than 1.0%, vanadium (V): less than 0.3%, titanium (Ti): less than 0.06%, niobium (Nb): less than 0.06%, and boron (B): less than 0.01%. The effect of these elements is saturated when these elements are added in the upper limit When the above elements are added in an amount exceeding the upper limit, the cost is increased. Therefore, when these elements are added, the amounts of the elements added should be not more than the above respective upper limits.

When phosphorus (P), sulfur (S), nitrogen (N), oxygen (O) and other elements, which are unavoidably included in general steels, are contained as steel constituents, the effect of the present invention is not deteriorated at all.

Further, in the coated steel sheet according to the present invention, in addition to the above elements and unavoidable impurities, elements, which are generally incidentally present in steels, may be contained as incidental constituents in such an amount that does not sacrifice the properties of the coated steel sheet.

The ductility of the steel sheet of the present invention as the final product varies depending upon the volume fraction of retained austenite contained in the product. The retained austenite contained in the metallurgical structure is stably present when the steel sheet does not undergo any deformation. Upon the application of deformation, however, the retained austenite

to

martensite

to

is

transformed

transformation induced plasticity. Therefore, in the steel sheet containing retained austenite in the metallurgical structure, good formability can be realized while enjoying high strength.

When the volume fraction of retained austenite is less than 2%, the above effect is not significantly attained. On the other hand, when the volume fraction of retained austenite exceeds 20%, there is a possibility that forming under extremely severe conditions provides a press formed product containing a large amount of martensite. As a result, problems associated with secondary formability and impact resistance sometimes occur. For the above reason, in the present invention, the volume fraction of retained austenite is limited to not more than 20%.

The ductility of the steel sheet of the present invention as the final product varies depending upon the volume fraction of retained austenite contained in the steel sheet as the final product. The retained austenite remaining in the metallurgical structure is stably present when the steel sheet does not undergo any deformation. Upon the application of deformation, however, the retained austenite is transformed to martensite to develop transformation induced plasticity. Therefore, good formability can be realized while enjoying high strength.

When the volume fraction of retained austenite is less than 2%, the effect of improving the formability is not significant. On the other hand, when the volume fraction of retained austenite exceeds 20%, there is a possibility that forming under extremely severe conditions provides a formed product containing a large amount of martensite. The presence of the martensite sometimes causes problems associated with secondary formability and impact resistance. For the above reason, in the present invention, the volume fraction of retained austenite is limited to not more than 20%.

35 (b) Zinc coated layer

5

10

15

20

25

30

In the steel sheet according to the present invention,

a zinc coated layer is provided on the steel sheet substrate. The zinc coated layer according to the present invention may be either a galvanized layer or a galvannealed layer. The galvanized layer and the galvannealed layer will be described in detail.

The galvanized layer comprises zinc: not less than 80% and aluminum: not more than 1% with the balance consisting of zincandunavoidable impurities. The reason why the content of zinc in the galvanized layer is limited to not less than 80% is that, when the zinc content is less than 80%, the coated layer is hard and is disadvantageously cracked at the time of forming. The reason why the content of aluminum in the galvanized layer is limited to not more than 1% is that, when the aluminum content exceeds 1%, aluminum segregated during coating constitutes a local battery which deteriorates corrosion resistance.

The galvannealed layer is useful particularly for improving spot weldability. The galvannealed layer comprises zinc: 80 to 91%, iron: 8 to 15%, and aluminum: not more than 1% with the balance consisting of zinc and unavoidable impurities. The reason why the content of zinc in the galvannealed layer is limited to not less than 80% is that, when the zinc content is less than 80%, the coated layer is hard and is disadvantageously cracked at the time of forming. The reason why the upper limit of the content of zinc in the galvannealed layer is 91% is that, when the zinc content exceeds 91%, the spot weldability is disadvantageously deteriorated making it impossible to attain the object of the present invention.

The reason why the content of iron in the galvaneealed layer is limited to not less than 8% is that, when the iron content is less than 8%, the suitability for conversion treatment (phosphate treatment) and the plating adhesion cannot be ensured. The reason why the upper limit of the content of iron in the coating layer is 15% is that, when the iron content exceeds 15%, overalloying occurs and,

consequently, plating adhesion in the worked part is deteriorated.

The reason why the content of aluminum in the galvannealed layer is limited to not more than 1% is that, when the aluminum content exceeds 1%, aluminum segregated during coating constitutes a local battery and, consequently, the corrosion resistance of the steel sheet is deteriorated.

The galvanized layer and the galvannealed layer in the steel sheet according to the present invention are as described above. In addition to the above constituents, elements such as manganese (Mn), lead (Pb), antimony (Sb), calcium (Ca), and magnesium (Mg) may be contained as unavoidable impurities. Further, very small amounts of other elements may be contained as incidental constituents.

The thickness of the galvanized layer and the galvannealed layer is not particularly limited. Preferably, however, the thickness is not less than 0.1 µm from the viewpoint of ensuring corrosion resistance, and is not more than 15 µm from the viewpoint of ensuring workability.

Production process

5

10

20

25

30

Next, the process for producing the hot-dip galvanized or galvannealed steel sheet according to the present invention will be described.

The hot-dip galvanized steel sheet according to the present invention is produced by annealing a cold rolled steel sheet having the above-described chemical composition for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, optionally further holding the cooled steel sheet in said cooling temperature range for not more than 10 min, then subjecting the cooled steel sheet to hot-dip galvanizing, and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

35 The hot-dip galvannealed steel sheet according to the present invention is produced by annealing a cold rolled steel

sheet having the above-described chemical composition for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, optionally further holding the cooled steel sheet in said cooling temperature range for not more than 10 min, then subjecting the cooled steel sheet to hot-dip galvannealing process, holding the coated steel sheet in the temperature region of 450 to 600°C for 5 sec to 2 min, and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

In continuous annealing of the cold rolled steel sheet after cold rolling, the cold rolled steel sheet is first heated to the temperature range of Ac₁ transformation point to Ac₃ transformation point to form a two-phase structure of [ferrite + austenite]. At that time, when the heating temperature is below 650°C, a lot of time is required for the redissolution of cementite to form a solid solution, and the existing amount of austenite is very small. For this reason, the lower limit of the heating temperature is 650°C.

10

15

20

25

30

35

On the other hand, when the heating temperature is excessively high, the volume fraction of austenite is so large that the content of carbon in austenite is lowered. For the above reason, the upper limit of the heating temperature is 900°C. When the holding time in this temperature range is excessively short, the possibility of presence of undissolved carbides is high and, consequently, the existing amount of austenite is small. On the other hand, when the holding time is long, grains become coarse and, as a result, the amount of austenite, which is finally present, is reduced, resulting in deteriorated strength-ductility balance. For the above reason, in the present invention, the holding time is limited to 10 sec to 6 min.

After the equation of heat, the steel sheet is cooled to 350 to 500°C at a cooling rate of 2 to 200°C/sec. The object of this step is as follows. In this step, austenite formed by heating in the two-phase region is carried forward to a

bainite transformation region without transformation to pearlite, and subsequent treatment permits retained austenite and bainite to exist at room temperature, whereby predetermined properties are provided. In this case, when the cooling rate is less than 2°C/sec, a major part of austenite disadvantageously causes pearlite transformation during cooling. As a result, retained austenite cannot be ensured. On the other hand, when the cooling rate exceeds 200°C/sec, the cooling termination temperature significantly deviates from a predetermined value in the widthwise direction and longitudinal direction. This makes it impossible to produce a steel sheet having homogeneous quality.

10

15

20

25

30

35

The termination temperature of cooling from the two-phase region is determined from the viewpoint of the suitability for hot-dip galvanizing. When the hot-dip galvanizing temperature is excessively low, the wettability of the steel sheet by coating is lowered and, consequently, plating adhesion is deteriorated. On the other hand, when the hot-dip galvanizing temperature is excessively high, an alloying reaction of iron with zinc proceeds in a zinc bath and, consequently, the concentration of iron in the coating is increased. For the above reason, in the present invention, the termination temperature of cooling from the two-phase region and the hot-dip zinc coating temperature are limited to 350 to 500°C.

Further, before hot-dip galvanizing, if necessary, the steel sheet is held in the temperature range of 350 to 500°C for not more than 10 min. Holding the temperature of the steel sheet before hot-dip galvanizing allows bainite transformation to proceed, and carbon-enriched retained austenite can be stabilized. As a result, steel sheets having a combination of good strength with good elongation can be more stably produced.

When the termination temperature of cooling from the two-phase region exceeds 500°C, austenite is decomposed to carbides during subsequent temperature holding, and it is

difficult for austenite to remain undecomposed. On the other hand, when the cooling termination temperature is below 350°C, a major part of austenite is transformed to martensite. As a result, the press formability is likely to be deteriorated, although the strength is enhanced. Further, in this case, at the time of galvanizing, the steel sheet temperature should be raised. This is ineffective from the viewpoint of thermal energy.

For the above reason, the holding temperature is limited to 350 to 500°C. When the temperature holding time exceeds 10 min, upon heating after zinc coating, the precipitation of carbides and the disappearance of untransformed austenite take place. As a result, both the strength and the press formability are likely to be deteriorated. For the above reason, when the temperature is held, the temperature holding time is limited to not more than 10 min.

10

15

20

25

30

35

In producing the hot-dip galvanized steel sheet, after hot-dip galvanizing process, the coated steel sheet is cooled to 250°C or below at a cooling rate of not less than 5°C/sec. Here at the time of galvanizing, bainite transformation is allowed to proceed to develop a mixed structure. The mixed structure comprises bainite, which is substantially free from carbides, retained austenite, which has been enriched with carbon scavenged from that portion and has an Ms point lowered to room temperature or below, and ferrite, which has been further cleaned during heating in the two-phase region. This structure can simultaneously realize high strength and good formability.

In this case, when the cooling rate after temperature holding is less than 5°C/sec or when the cooling termination temperature is above 250°C, austenite, which has been enriched with carbon during cooling, also causes carbide precipitation and is decomposed to bainite. This disadvantageously reduces the amount of retained austenite which improves workability through transformation induced plasticity. As a result, the object of the present invention cannot be attained. In order

5

10

15

20

25

30

35

to increase the amount of the retained austenite remaining untransformed, preferably, the holding temperature after hot-dip galvanizing process is 350 to 400°C, and the holding time is not more than 5 min.

Further, in producing the hot-dip galvannealed steel sheet, after hot-dip galvannealing process, the coated steel sheet is held in the temperature range of 450 to 600°C for 5 sec to 2 min and is then cooled to 250°C or below at a cooling rate of not less than 5°C/sec. The above conditions are determined from the viewpoints of the alloying reaction of iron with zinc and the optimization of the structure of the steel sheet.

In the steel according to the present invention, silicon and aluminum are contained, and through the utilization of two-stage transformation from austenite to bainite, a mixed structure is developed which is composed of bainite, retained austenite, and ferrite. This bainite is substantially free from carbides. The austenite has been enriched with carbon scavenged from that portion and has an Ms point lowered to room temperature or below. The ferrite has been further cleaned during heating in the two-phase region. development of the mixed structure can simultaneously realize high strength and good formability. When the holding temperature is above 600°C, pearlite is formed and the retained austenite is not contained in the steel sheet. Further, in this case, the alloying reaction excessively proceeds. Consequently, the concentration of iron in the coating disadvantageously exceeds 12%.

On the other hand, when the heating temperature is 450° C or below, the alloying reaction rate of the coating is lowered and, consequently, the concentration of iron in the coating is lowered.

When the holding time is not more than 5 sec, bainite is not satisfactorily formed and the enrichment of untransformed austenite with carbon is also unsatisfactory. In this case, during cooling, martensite is formed, resulting

5

10

15

25

30

35

in deteriorated formability. At the same time, the galvannealing reaction is unsatisfactory.

On the other hand, when the holding time is not less than 2 min, overalloying of the coating occurs and, at the time of forming, coating separation or the like is disadvantageously likely to occur. When the cooling rate after the temperature holding is less than 5°C/sec or when the cooling termination temperature is above 250°C, bainite transformation further proceeds and austenite, which has been enriched with carbon in the preceding reaction, also causes carbide precipitation and is decomposed to bainite. This disadvantageously reduces the amount of retained austenite which improves workability through transformation induced plasticity. As a result, the object of the present invention cannot be attained.

The hot-dip galvanizing temperature is preferably between the melting point of the zinc bath and 500°C. When the hot-dip galvanizing temperature is above 500°C, a large amount of vapor is produced from the zinc bath and, consequently, the operating efficiency is deteriorated. The rate of heating to the holding temperature after the coating is not particularly limited. The heating rate, however, is preferably not less than 3°C/sec from the viewpoints of the coating structure and the metallurgical structure.

The temperature and the cooling temperature in the above-described individual steps are not necessarily constant so far as the temperature and the cooling temperature fall within the above-specified respective ranges. Even when the temperature or the cooling temperature fluctuates within the above-specified range, the properties of the final product are not deteriorated and, in some cases, are improved. The material used in the present invention may have been produced through refining, casting, hot rolling, and cold rolling steps in a conventional steelmaking process. Alternatively, the material used in the present invention may have been produced by a process wherein a part or the whole of these steps has

been omitted. Conditions of these steps are also not particularly limited.

In order to further improve the plating adhesion, before annealing, the steel sheet may be coated with nickel, copper, cobalt, and iron, either alone or in combination. Another method usable for improving the plating adhesion is to properly regulate the atmosphere at the time of annealing of the steel sheet. For example, a method may be adopted wherein, before coating, the surface of the steel sheet is first oxidized in atmosphere and is then reduced to clean the surface of the steel sheet. Further, for plating adhesion improvement purposes, before annealing, pickling of the steel sheet or grinding of the steel sheet may be carried out to remove oxides on the surface of the steel sheet. This does not change the subject matter of the present invention. The above treatments can improve plating adhesion and further can accelerate alloying.

As described above, the present invention can efficiently produce high strength hot-dip galvanized or galvannealed steel sheets having good press formability and plating adhesion which can be used as automobile, building, electric or other members and other applications.

[EXAMPLES]

The following examples further illustrate but do not limit the present invention.

Example A1

5

10

15

20

30

35

Steels having chemical compositions shown in Table Al were reheated to 1250°C, were then finish rolled at 900°C, and were coiled at 650°C to prepare 4 mm-thick hot rolled steel sheets. Scale deposited on the surface of the hot rolled steel sheets was removed with hydrochloricacid, and the steel sheets were then cold rolled to prepare 1.4 mm-thick cold rolled steel sheets. The coldrolled steel sheets were annealed under conditions shown in Table A2 and Table A3 (continuation from Table A2), were coated, and were then temper rolled with a

reduction of 0.5%. The steel sheets thus obtained were subjected to the following performance evaluation tests, that is, "tensile test," "retained austenite measuring test," "welding test," "appearance of coating," "plating adhesion," and "measurement of concentration in coated layer." In the coating in the above production of the steel sheets, both sides of the cold rolled steel sheets were coated at a coverage of coating of 50 g/m^2 per side.

5

10

15

20

25

30

35

In the "tensile test," a JIS No. 5 tensile test piece was extracted in C-direction, and a cold tensile test was carried out under conditions of gauge thickness 50 mm and tensile speed 10 mm/min.

The "retained austenite measuring test" was carried out by a method called "5-peak" method. In this method, a quarter of the sheet thickness from the surface toward the inner side of the sheet was chemically polished, α -iron intensity and γ -iron intensity were then measured by X-ray diffractometry using an Mo bulb, and the volume fraction of retained austenite was determined based on the α -iron intensity and the γ -iron intensity.

The "welding test" was carried out by performing spot welding under welding conditions of welding current: 10 kA, applied pressure: 220 kg, welding time: 12 cycles, electrode diameter: 6 mm, electrode shape: domed, and tip: 6ϕ -40R, and counting the number of continuous spots provided until the welding reached the point at which the nugget diameter became below $4\sqrt{t}$ wherein trepresents sheet thickness. The counted number of continuous spots was evaluated according to the following criteria. O: more than 1,000 continuous spots, Δ : 500 to 1,000 continuous spots, and \times : less than 500 continuous spots. Here O was regarded as acceptable, and Δ and \times as unacceptable.

The "appearance of coating" was determined by visually inspecting the appearance of the coated steel sheet for noncoated sites and evaluating the results according to the following criteria. \bigcirc : not more than $5/dm^2$, \triangle : 6 to $15/dm^2$,

and \times : notless than 16/dm². Here \bigcirc was regarded as acceptable, and \triangle and \times as unacceptable.

The "plating adhesion" was determined by subjecting the coated steel sheet to a 60-degree V bending test, then performing a tape test, and evaluating the results according to the following criteria.

Blackening in tape test (%)

©: 0 to 10

10

15

20

25

30

35

O: 10 to less than 20

 \triangle : 20 to less than 30

X: not less than 30

(\bigcirc and \bigcirc : acceptable, \triangle and \times : unacceptable)

The "measurement of concentration in coated layer" was carried out by dissolving the coated layer in 5% hydrochloric acid containing an amine-based inhibitor and then analyzing the solution by ICP emission spectroscopy.

The results of the above performance evaluation tests are shown in Table A4 and Table A5 (continuation from Table A4). For all of samples 1 to 13 which are examples of the present invention, the total elongation was not less than 30% while ensuring a tensile strength of not less than 550 MPa, indicating that high strength and good press formability could be simultaneously realized. At the same time, for these samples, the plating adhesion was acceptable.

By contrast, samples 14 to 23, which are comparative examples, could not attain the object of the present invention, because, for sample 14, the content of carbon was lower than the carbon content range specified in the present invention; for sample 15, the content of carbon was higher than the carbon content range specified in the present invention; for sample 16, the content of silicon was lower than the silicon content range specified in the present invention; for sample 17, the content of silicon was higher than the silicon content range specified in the present invention; samples 18 and 19 failed to satisfy the relationship between silicon and aluminum specified in the present invention; for sample 20, the content

of manganese was lower than the manganese content range specified in the present invention; for sample 21, the content of manganese was higher than the manganese content range specified in the present invention; for sample 22, the content of aluminum was higher than the aluminum content range specified in the present invention; and, for sample 23, the content of tin was lower than the tin content range specified in the present invention.

5

10

15

For samples 24 to 48 as comparative examples, although the chemical compositions of the steels fell within the scope of the present invention, any one of the treatment conditions was outside the scope of the present invention. These comparative samples suffered from poor strength-ductility balance or poor plating adhesion and thus could not attain the object of the present invention.

		ŋ	n.	n.	inc.	7	2	Š	ž	>	2	Inv.	lnv.	120	2	Γ									
		Remarks	. of inv.	1	Å	9	1			•	1	8	of	of inc	of 1mv	Colle	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.	Comp. Ex.
	L	~	EX	EX.	X	Ä	BX.	EX	EX.	EX.	ß.	EX.	EX.	E E	MX.	ပိ	8	8	8	8	8	8	8	8	8
		Si+0.8Al	1.41	0.824	1.316	1.242	1.468	0.86	1.884	1.092	1.22	1.136	1.042	0.924	1.584	0.78	0.796	0.442	2.112	0.316	2.134	0.536	1.162	1.844	0.876
	-	Others	1	Mo: 0.11, Ca: 0.02	Cr: 0.22	sb: 0.12	Ce (REM): 0.002, Nb: 0.03	Ti: 0.02, Zr: 0.05	Mg: 0.003	ı	X (REM): 0.07, Ca: 0.01	1	Mo: 0.04, Ti: 0.01, Mg: 0.02	1	Mg: 0.02	1	Bi: 0.05	Ce (REM): 0.2	1	V: 0.01, Zr: 0.02	1	Sb: 0.22, Ca: 0.21	Cr: 0.23, Mo: 0.09	ŀ	Ti: 0.03, Nb: 0.03
	weight%	တိ		0.02	ı	1.	•	0.23	ı	ı	,	0.01	0.01	0.02	!		\$	0.12	1	0.04	ı	1	1	0.12	,
3 A1	on, we	Cu	0.01	0.04	0.20	1.30	0.02	0.20	0.10	0.92	0.02	0.65	90.0	0.01	0.10	0.03	0.02	0.02	0.10	0.23	0.13	0.22	0.15	1.10	0.03
Table Al	positi	'nŢ	0.02	0.05	0.03	0.87	90.0	1.60	0.20	0.67	0.11	1.12	0.23	0.02	0.20	0.01	0.02	0.02	1.20	0.02	0.08	0.66	0.23	0.01	0.32
	Chemical composition,	Sn	0.056	0.015	0.720	0.132	0.082	0.008	0.432	0.187	0.861	0.006	0.229	0.046	0.079	0.028	0.102	0.043	0.081	0.210	0.192	0.062	0.009	0.008	0.001
	Chem	Al	0.25	0.73	0.02	0.74	0.21	0.65	0.23	0.34	0.95	0.02	0.14	0.13	0.23	0.55	0.22	0.39	0.24	*0.12	*0.73	0.22	0.54	1.63	0.32
		S	0.005	0.003	0.004	0.006	0.002	0.012	0.008	0.004	0.011	0.005	0.009	0.005	0.008	0.005	0.002	0.003	0.002	0.007	0.003	0.003	0.005	0.006	0.002
		ρı	0.004	0.014	0.008	0.00	0.015	0.012	0.005	0.005	0.012	0.004	0.008	0.004	0.005	0.003	0.013	0.007	0.008	0.004	0.005	0.008	0.009	0.002	0.012
		Mn	1.55	2.21	1.40	1.00	2.40	0.00	1.50	1.40	1.60	1.30	1.6	1.70	1.70	1.03	1.82	1.34	0.97	0.58	1.52	0.18	2.65	0.85	1.22
		Si	1.21	0.24	1.30	0.65	1.30	0.34	1.70	0.82	0.46	1.12	0.93	0.82	1.40	0.34	0.62	0.13	1.92	*0.22	*1.55	0.36	0.73	0.54	0.62
		υ	0.08	0.12	0.16	0.13	0.04	0.07	0.17	0.09	0.11	0.07	0.18	0.08	0.17	0.01	0.22	0.13	0.16	0.15	0.12	90.0	0.14	0.12	0.09
	Stopl		ъ	q	υ	ס	9	£	ה	Д	7		א	1	Ħ	ជ	٥	Q	54	н	80	44	p	>	3

Note) Mark ____indicates that the value is outside the scope of the present invention. Mark * indicates that the value does not satisfy the relationship between Al and Si specified in the present invention. Mark — indicates that the constituent is not contained.

Remarks ñ B G G ð ð ß G ð ð ક క G ð g Cooling temp., Cooling ್ಗೆ/3ec rate, ដ 위 위 Alloying time, Alloying temp., ပူ Al content of zinc 0.15 0.11 0.14 0.16 0.10 0.12 0.15 0.20 0.11 0.15 0.20 0.18 0.20 0.15 bath, & 0.14 0.09 0.16 0.13 0.10 0.18 0.17 0.09 0.14 0.08 Coating temp., ပူ Holding time, Table A2 Sec temp. before ပူ - 400 400 - 450 360 - 440 - 500 360 - 440 400 - 470 - 490 - 470 400 - 470 370 - 440 coating, ı termination Cooling temp. Cooling °C/sec rate, ដ m ^ Ŋ time, sec Annealing Annealing temp., °C Steel Д U v Ø Þ × E Ď, Ø ų Sample ~ m 2 2 œ

indicates that the value is outside the scope of the present invention. The heating rate after coating was constant and 10°C/sec. GA represents a hot-dip galvannealed steel sheet. GI represents a hot-dip galvanized steel sheet. Note Mark

Table A3 (continuation from Table A2)

				Cooling	Cooling	Holding	Holding	Coating	Al content	Alloying	Alloying	Cooling	Cooling	
Sample	Steel	Anneali		rate,	termination		time,	temp.,	of zinc	temp.,	time,	rate,	temp.,	Remarks
		temp., 'C	time, sec	°C/sec	temp.,°C	coating, "C	sec	ပ္	bath, &	ပ္စ	sec	°C/sec	ပ္	
25	п	950	06	08	480	400 - 500	09	450	0.11	480	50	10	180	ક
26	В	750	S	20	440	430 - 500	20	450	0.20	500	30	10	180	ક
27	ø	800	400	20	410	-	•	480	0.17	500	40	20	220	g.
82	B	750	06	1	370	430 - 500	30	450	0.13	510	30	10	180	8
53	rđ	800	110	10	008	370 - 440	300	450	60.0	480	50	10	180	ક્
30	ď	077 /	06	70	520	370 - 440	300	450	0.14	500	30	10	180	æ
31	r	830	150	10	420	400 - 490	650	480	0.18	500	30	10	180	g.
32	rs	800	70	20	410	400 - 470	80	430	0.11	430	40	15	180	G.A.
33	В	750	06	25	440	370 - 440	140	480	0.16	620	20	10	100	G.
34	đ	850	09	20	420	1	•	450	0.20	500	m	80	180	કુ
35	æ	750	90	80	480	•	-	450	0.12	500	130	10	180	g.
36	ď	820	70	50	490	400 - 470	250	440	0.10	500	25	6	180	કુ
37	ď	750	100	20	360	1	•	450	0.08	500	30	10	300	કુ
38	В	830	06	20	480	400 ~ 500	09	450	0.82	500	25	. 10	180	g.
39	rs	009	06	20	360	_	•	470	0.08	•	•	10	180	ĞI
40	a	950	90	80	480	400 - 500	60	450	0.11	8	•	10	180	GI
41	æ	750	5	20	440	430 - 500	20	450	0.20	•	•	10	180	19
42	R	800	420	20	410	1	•	480	0.17	ı	•	20	220	ij
43	B	750	90	1	370	430 - 500	30	450	0.13		•	10	180	GI
44	ď	800	110	10	300	370 - 440	300	450	0.09	•	1	10	180	ID
45	ъ	830	150	10	420	400 - 490	720	480	0.18		•	10	180	GI
46	æ	820	70	50	490	400 - 470	250	440	0.10	1	•	m	180	ß
47	r	750	. 100	20	360	•	•	450	0.08	•	•	ខ្ម	300	19
48	В	830	06	20	480	400 - 500	09	450	0.82	•		10	180	GI

Note) Mark indicates that the value is outside the scope of the present invention. The heating rate after coating was constant and 10°C/sec. GA represents a hot-dip galvannealed steel sheet. GI represents a hot-dip galvanized steel sheet.

Table A4

		d	Ketained	Fe content of	Al content of	Appearance	Plating		
•	43, FEG		ነ, ቄ	coating, 8	coating, 8	of coating	adhesion	Weldability	Kemarks
-	630	98	8	01	0.22	Ö	0	0	Ex. of inv.
2	099	34	10	10	0.18	0	0	0	Ex. of inv.
9	720	34	13	-	0.28	0	0	0	Ex. of inv.
9	640	31	11	11	0.16	0	0	0	Ex. of inv.
S	780	30	က		0.18	0	0	0	Ex. of inv.
9	580	36	6	10	0.82	0	0	0	Ex. of inv.
7	840	31	15	10	0.13	0	0	0	of
8	640	36	8	12	0.16	0	0	0	Ex. of inv.
6	630	35	10	•	0.25	0	0	0	1
21	620	36	7	10	0.17	0	0	0	Ex. of inv.
11	810	31	18	10	0.23	0	0	0	Ex. of inv.
12	610	35	4	11	0.32	0	0	0	ı
13	830	32	15	1	0.18	0	0	0	Ex. of inv.
14	540	28	7	ŀ	0.20	0	0	0	Comp. Ex.
15	810	25	22	10	0.21	0	0	×	Comp. Ex.
16	570	28	1	10	0.16	0	0	0	Comp. Ex.
17	710	30	8	9	0.32	×	×I	0	Comp. Ex.
18	550	29	1	10	0.13	0	· (0	0	Comp. Ex.
19	620	33	9	10	0.19	×	×I	0	Comp. Ex.
20	560	20	1	6	0.23	0	0	0	Comp. Ex.
21	700	34	7	10	0.28	Q	⊲	×	Comp. Ex.
22	640	33	9	1	0.32	×[×I	0	Comp. Ex.
23	099	33	S	3	0.28	×	×	0	Comp. Ex.
24	550	24	1	10	0.41	0	0	0	Comp. Ex.

Table A5 (continuation from Table A4)

				- i	TOIL TADIE A4)			
MPa	E1, 8	Retained 7, 8	Fe content of coating, 8	Al content of coating, &	Appearance of	Plating	Weldability	Remarks
009	26	1	10	1-4	0	©	С	Comp. Ex
620	20	-1	6	0.23	С	©	C	Comp Ev
580	22	1	10	0.63	0	@	C	Comp. Ex
550	56	1	10	0.27	0	0	C	Comp
650	24	1	11	0.34	0	0	C	Comp. Ex
610	34	9	15	0.28	0	□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	0	Comp. Ex.
009	29	1	10	0.41	0	0	0	Comp. Ex.
610	35	5	5	0.23	0	0	0	Comp. Ex.
570	29	1	15	0.23	0	٥	0	Comp. Ex.
630	35	9	Ī	0.23	0	0	0	Comp. Ex.
580	28	F	15	0.32	0	V	0	Comp. Ex.
580	26	1	10	0.23	0	0	0	Comp. Ex.
560	23	1	10	0.32	0	0	0	Comp. Ex.
630	35	7	10	1.23	0	0	0	Comp. Ex.
550	24	1	10	0.41	0	0	0	Comp. Ex.
009	26	1	10	0.21	0	0	0	Comp. Ex.
620	20	1	6	0.23	0	0	0	Comp. Ex.
580	22	1	10	0.63	0	0	0	Comp. Ex.
550	26	1	10	0.27	0	0	0	Comp. Ex.
650	24	1	11	0.34	0	0	0	Comp. Ex.
009	29	1	10	0.41	0	0	0	Como. Ex.
580	26	1	10	0.23	0	0	0	Comp. Ex.
560	23	1	10	0.32	0	0	С	Comp. Ex.
630	35	7	10	1.23	0	0	С	Comp. Ex

Note) Mark ____ indicates that the value is outside the scope of the present invention.

Example B1

5

10

15

20

25

30

35

Steels having chemical compositions shown in Table B1 and Table B2 (continuation from Table B1) were reheated to 1250°C, were then finish rolled at 900°C, and were coiled at 650°C to prepare 4 mm-thick hot rolled steel sheets. deposited on the surface of the hot rolled steel sheets was removed with hydrochloric acid, and the steel sheets were then cold rolled to prepare 1.4 mm-thick cold rolled steel sheets. The cold rolled steel sheets were annealed under conditions shown in Table B3 and Table B4 (continuation from Table B3), were coated, and were then temper rolled with a reduction of 0.5%. The steel sheets thus obtained were subjected to the following performance evaluation tests, that is, "tensile test," "retained austenite measuring test," "welding test," "appearance of coating," "plating adhesion," and "measurement of concentration in coated layer." In the coating in the above production of the steel sheets, both sides of the cold rolled steel sheets were coated at a coverage of coating of 50 g/m^2 per side.

In the "tensile test," a JIS No. 5 tensile test piece was extracted in C-direction, and a cold tensile test was carried out under conditions of gauge thickness 50 mm and tensile speed 10 mm/min.

The "retained austenite measuring test" was carried out by a method called "5-peak" method. In this method, a quarter of the sheet thickness from the surface toward the inner side of the sheet was chemically polished, α -iron intensity and γ -iron intensity were then measured by X-ray diffractometry using an Mo bulb, and the volume fraction of retained austenite was determined based on the α -iron intensity and the γ -iron intensity.

The "welding test" was carried out by performing spot welding under welding conditions of welding current: 10 kA, applied pressure: 220 kg, welding time: 12 cycles, electrode diameter: 6 mm, electrode shape: domed, and tip: 6ϕ -40R, and counting the number of continuous spots provided until the

welding reached the point at which the nugget diameter became below $4\sqrt{t}$ wherein t represents sheet thickness. The counted number of continuous spots was evaluated according to the following criteria. \odot : more than 2,000 continuous spots, \odot : more than 1,000 continuous spots, Δ : 500 to 1,000 continuous spots, and \times : less than 500 continuous spots. Here \odot and \odot were regarded as acceptable, and Δ and \times as unacceptable.

The "appearance of coating" was determined by visually inspecting the appearance of the coated steel sheet for non-coated sites and evaluating the results according to the following criteria.

 \bigcirc : not more than $5/dm^2$, \triangle : 6 to $15/dm^2$, and \times : not less than $16/dm^2$.

15 Here \bigcirc was regarded as acceptable, and \triangle and \times as unacceptable.

The "plating adhesion" was determined by subjecting the coated steel sheet to a 60-degree V bending test, then performing a tape test, and evaluating the results according to the following criteria.

Blackening in tape test (%)

©: 0 to 10

5

10

20

25

30

35

O: 10 to less than 20

 \triangle : 20 to less than 30

X: not less than 30

(\bigcirc and \bigcirc : acceptable, \triangle and \times : unacceptable)

The "measurement of concentration in coated layer" was carried out by dissolving the coated layer in 5% hydrochloric acid containing an amine-based inhibitor and then analyzing the solution by ICP emission spectroscopy.

The results of the above performance evaluation tests are shown in Table B5 and Table B6 (continuation from Table B5). For all of samples 1 to 13 which are examples of the present invention, the total elongation was not less than 30% while ensuring a tensile strength of not less than 550 MPa, indicating that high strength and good press formability

5

10

15

20

25

30

35

could be simultaneously realized. At the same time, for these samples, the plating adhesion was acceptable.

By contrast, samples 14 to 26, which are comparative examples, could not attain the object of the present invention due to poor strength-ductility balance or poor plating adhesion, because, for sample 14, the content of carbon (C) was lower than the carbon (C) content range specified in the present invention; for sample 15, the content of carbon (C) was higher than the carbon (C) content range specified in the present invention; for sample 16, the content of silicon (Si) was lower than the silicon (Si) content range specified in the present invention; for sample 17, the content of silicon (Si) was higher than the silicon (Si) content range specified in the present invention; samples 18 and 19 failed to satisfy the relationship between silicon (Si) and aluminum (Al) specified in the present invention; for sample 20, the content of manganese (Mn) was lower than the manganese (Mn) content range specified in the present invention; for sample 21, the content of manganese (Mn) was higher than the manganese (Mn) content range specified in the present invention; for sample 22, the content of aluminum (Al) was higher than the aluminum (Al) content range specified in the present invention; for samples 23, 24, and 25, the content of selenium (Se) + bismuth (Bi) + antimony (Sb) was lower than the selenium + bismuth + antimony content range specified in the present invention; and, for sample 26, the content of selenium (Se) + bismuth (Bi) + antimony (Sb) was higher than the selenium + bismuth + antimony content range specified in the present invention.

For samples 27 to 51 (see Table B6) as comparative examples, although the chemical compositions of the steel sheets fell within the scope of the present invention, any one of the treatment conditions was outside the scope of the present invention. These comparative samples suffered from poor strength-ductility balance or poor plating adhesion and thus could not attain the object of the present invention.

Table B1

									100 91	i ch th			Domarka	
						ບ	Chemical composition, weighte	Compost	TOTT'	- American		CALO RAI	CVTOMEN	_
Steel				6	6	7	ų,	Bi	Se	Sb+Bi+Se	Others	STTO. OUT		Т
	υ	21	Mn	4	2	1	3	1		0.056		1.41	Ex. of Inv.	<u>.</u>
٦	0.08	1.21	1.55	0.004	0.005	0.25	0.056		2	20.0	7 7 7 7 7 7	0 824	Ex. of inv.	>
,	5	1	2 21	0.014	0.003	0.73	0	0.050	0	0.050	MO: U.11, CA: U.02		The American	T:
٩	21.0		ı		100	20	c	c	0.022	0.022	sn: 0.05, Cr: 0.22	1.310	EA. OL AN	;
ช	0.16	1.30	1.40	0.008	0.004	0.02	200	000	120	0 482	Ni: 0.12	1.242	Ex. of inv.	[خ
7	0.13	0.65	1.00	0.009	0.006	0.74	0.132	0.230	77.0	205.0	Co. /BEM . 0 002 Nb: 0.03	1.468	Ex. of inv.	>
	0 04	1 30	2.40	0.015	0.002	0.21	0.082	0.060	0.020	0.102	יייי ייייי אייייי פייייי	78 0	Ex of inv	5
,				0 013	0 012	0.65	0.008	0.002	100.0	0.011	Ti: 0.02, Zr: 0.05	20.0	; Y	1
н	0.01	0.34	30	0.012	7:01		5	0000	100	0 412	Mg: 0.003	1.884	EX. OF THE	<u>.</u>
ь	0.17	1.70	1.50	0.002	0.008	0.23	0.112	0.400	207.0			1.092	Ex. of inv.	>.
,	300	500	,	0 005	0 004	0.34	0.187	0	0.030	177.0			١	:
ч	0.09	0.09 0.82	7.40	3			27.0	0110	0.00	0 872	Y (REM): 0.07, Ca: 0.01	1.22	EX. OF THY	
٠,٦	0.11	0.46	1.60	0.012	0.011	0.95	0./42	0.110	0.020	20.0		1.136	Ex. of inv.	₹.
	100		1 30	0.004	0.005	0.02	900.0	0.320	0	0.326				Τ
			+						0	0110	Mo: 0.04, Ti: 0.01,	1.042	Ex. of inv.	<u>.</u>
	0.18	0.93	٦.6	0.008	0.009	0.14	0.229	0.230 0.060	0.0	670.0	Mg: 0.02			7
:							3.0	000	010	0 076	ŧ	0.924	Ex. of inv.	<u>.</u>
7	0.08	0.82	1.70	0.004	0.005	0.13	0.046	0.020	2 .	\perp	Mar. 0 02 Co. 0 05	1.584	Ex. of inv.	
1	2.0	0 17 1 40	1 70	0.005	0.008	0.23	0	0.200	0.100	0.300	my. 0.02, 00: 0:0]
Ħ	17:0	2	4	4	4				- 4	30 0000	the process invention;			
						4 4 4	C		מכיי		CITE DIESCIT THE CITE TO			

Note) In the table, mark ___indicates that the value is outside the scope of the present invention; and mark * indicates that the value does not satisfy the relationship between Al and Si specified in the present invention; and mark -- indicates that the constituent is not contained.

Table B2 (continuation from Table B1)

							-						
Steel	ļ			•		Cne	mical co	mpositio	unemical composition, weights	را			
	اد	ST	Wh	<u>a</u>	S	Z	Sb	Bi	Se	Sb+Ri+Se	Others	LEG OFFE	Remarks
G	0.01	0.34	1.03	0 003	200	7.5	000	0,00	000	2000	STRIP	THO . OLTO	
•	33	5					0.70	0.010	0.030	0.058		0.78	COMD. EX.
}	77.	70.0	7.87	0.013	0.002	0.22	0.102	0.020	0.020	0.142	Sp: 0.05	0 796	1 Campo
۵,	0.13	0.13	1.34	0.007	0.003	0.39	0.043	0.020	0	690.0	Co (BEM) 0 2	26.0	- Table 1
ש	0.16	1.92	0.97	0.008	0 002	0 24	700	000	500	000	2.0 : (TENT) 30	7.4.0	Comp. EX.
٠	0 15	\$0.00	000	100	100		100.0	0.002	2005	0.008	,	2.112	Comp. Ex.
		33:2	0.30	700.0	0.00/	-0.12	0.210	0.020	0.230	0.460	V: 0.01, Zr: 0.02	0.316	Comp R.v
8	0.12	*1.55	1.52	0.002	0.003	*0.73	0.192	0.080	0.130	0 402		200	
t t	90.0	0.36	0.18	0.008	0 003	0 22	0 063	0,00		202.0		2.134	COMD. EX.
];	7					77.0	700.0	0.042	0.220	0.324	Cu: 0.22, Ca: 0.21	0.536	Comp. Ex.
3	7.14	0.73	2:05	0.00	0.005	0.54	0	0.230	0.150	0.380	Cr: 0.23 Mo: 0.09	1 163	- A - A - A - A - A - A - A - A - A - A
>	0.12	0.54	0.85	0.002	900.0	1.63	0.008	0.010	0.023	0 041		20111	Comp. Ex.
3	0.09	0.62	1.22	0.012	0.002	32	001		,			##0.1	Comp. EA.
*	000	69 0	1 22	3	000	3		3		0.00I	TI: 0.03, ND: 0.03	0.876	Comp. Ex.
		20.05	77.7	0.012	0.002	0.32	0	0.002		0.005	Ni: 0.11. Mg: 0.02	0.876	Comp De
Y	0.09	0.62	1.22	0.012	0.002	0.32	0.001	0	0.001	0.002	8n. 0 04	2000	- War - War - Co
N	0.09	0.62	1.22	0.012	0.002	0.32	0 861	0000	220		1 2.00	0.0.0	Comp. Ex.
]:						***	2.4%	0.447	7.730		0.876	Comp. Ex.

Note) In the table, mark ___indicates that the value is outside the scope of the present invention; mark * indicates that the value does not satisfy the relationship between Al and Si specified in the present invention; and mark - indicates that the constituent is not contained.

		10	_	_	$\overline{}$	_	\neg		_	_	\neg	7	Т	_		1	1	$\overline{}$	_	т	_	1		_	_	-	_		_	
		Remarks	6	5 6	5	5	45	5 6	3 6	\$	5	5	3	€	8	gi	15	4	8	89	8	8	8	5 8	5 5	3	3	8	ij	4
	Cooling	temp.,	,	160	200		087	087		001	180	180	180	180	180	200	180	180	180	180	180	180	180	180	180		200	180	180	180
	Cooling	rate,	1000	1.5	3 5	3 5		2 5	2 5	2 5		2	2	2	10	10	ខ្ព	2	ន	2	2	27	10	200	2	5		2	2	10
	Alloying	time,	35	ş	3 .	AC		' 	5	36	3	, ;	62	30	10	•	•	15 .	25	30	20	25	25	25			90	5	•	25
	ם ב	temp.,	500	520		200		1	005	2	3 '	000	200	200	550	1	ŧ	520	200	580	200	200	500	200		,	003	8	·	200
	Coating Al content of	zinc bath, &	0.11	0.14	0.16	0.10	13	0.15	0.15	0 20	0 11	9. 0		0.40	0.18	0.20	0.15	0.14	0.10	60.0	0.16	0.13	0.18	0.17	0.09	0.14	0 14		1	0.14
	Coating	temp., °q	450	450	470	450	440	450	450	450	450	450	98		450	460	450	460	450	450	450	450	450	450	450	440	440	440		0,4
Table B3	E	Sec.	09		420		300			6	200	5		1	0	'	8	-	,	30		300	•	30	90	ı		2	1	
Tab Holding	tom before	coating, °C		1	360 - 400	,	350 - 380	h		400 - 470	360 - 440	} •	١,	260 - 440	.	•	400 - 470	-	-	430 - 500	•	370 - 440	-	400 - 490	400 - 470	•		400 - 470	١,	1
Cooling	termination	temp., °C	400	400	420	380	370	480	410	440	360	480	440	908		430	410	420	370	480	410	440	420	480	490	400	400	400	440	
Cooling	rate	•	10	20	40	150	Э	20	20	7	10	20	s,	65		2	20	9	20	20	20	6	2	2	2	2	10	s	20	1
	Annealing	time, sec	150	90	100	90	150	06	200	85	180	06	110	120	200	3	2	120	110	90	06	150	70	90	200	120	110	120	70	1
	Anneal	temp.,°C	800	750	800	750	780	800	750	850	750	800	750	700	Bon	200	000	000	750	078	/30	0/0	00/	830	750	800	750	800	800	the table
	Steel	\rightarrow	В	ρ	U	ט	ø	¥	6	۾	7	î	×	τ	E		= (5	Δ,	0,	•	n .	١,	3	>	3	×	>	2	Ta th
	Sample		1	~	6	4	S	v	-	8	a	10	11	12	13	1	:		9 :	٩	2			7 8	3	2	24	25	26	Note)

GA represents a hot-dip galvannealed steel sheet, and GI represents a hot-dip galvanized steel sheet. In the table, mark indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and Note)

Table B4 (continuation from Table B3)

				Cooling	Cooling	Holding	Holding	Coating	Al content	Alloying	Alloying	Cooling	Cooling	
Sample	Steel	Anneali	Annealing	rate,	termination	temp. before	time,	temp.,	of zinc	temp.,	time,	rate,	temp.,	Remarks
		cemp., c	time, sec	°C/sec	temp.,°C	coating, °C	200	ပ္	bath, &	ပ္စ	sec	°C/sec	ပ္	
27	B	009	06	20	360	1	,	470	0.08	500	30	10	180	8
28	e	950	06	90	480	400 - 500	09	450	0.11	480	50	10	180	ક
29	æ	750	6 0)	20	440	430 - 500	20	450	0.20	200	30	10	180	8
30	В	800	400	20	410	-	-	480	0.17	200	40	20	220	G.
31	п	750	06	-1	370	430 - 500	30	450	61.0	510	30	10	180	G.
32	ъ	800	110	10	300	370 - 440	300	450	60.0	480	50	10.	180	8
33	a	770	06	70	. 220	370 - 440	300	450	0.14	500	30	10	180	đ
34	а	830	150	10	420	400 - 490	650	085	91.0	200	30	10	180	đ
35	а	800	70	20	410	400 - 470	80	430	0.11	430	40	15	180	প্ত
36	æ	750	06	25	440	370 - 440	140	480	0.16	620	20	10	100	æ
37	В	850	90	20	420	•	-	450	0.20	500	Э	8	180	45
38	В	750	06	80	480	•	-	450	0.12	500	130	10	180	GA
39	в	820	70	30	490	400 - 470	250	440	0.10	500	25	ю	180	প্ৰ
40	В	750	100	20	360	•	•	450	0.08	500	30	10	300	GA
41	В	830	06	20	480	400 - 500	60	450	0.82	200	25	10	180	GA
42	В	009	90	20	360	-	•	470	0.08	ı	,	10	180	GI
43	ď	950	06	80	480	400 - 500	60	450	0.11	•	,	10	180	ĞI
44	ъ	750	8	20	440	430 - 500	20	450	0.20	ţ	1	10	180	GI
45	a	800	420	20	410	•	-	480	0.17	•	•	20	220	GI
46	а	750	06	1	370	430 - 500	30	450	0.13	•	•	10	180	ΒĪ
47	æ	800	110	10	300	370 - 440	300	450	0.09	•	•	10	180	GI
48	В	088	150	10	420	400 - 490	720	480	0.18	ı	•	10	160	GI
49	æ	920	70	20	490	400 - 470	250	440	0.10	•	1	m	180	GI
20	a	750	100	20	360	•	-	450	0.08	,	•	10	300	GI
SI	æ	830	06	20	480	400 - 500	09	450	0.82	1	•	10	180	GI
Note)	III C	In the table, mark		indicates that	the value	is outside the	scope of		the present invention;	ë				

In the table, mark indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and GA represents a hot-dip galvanized steel sheet. Note)

Table B5

		Kemarks	Ex. of inv.	of.	of.	of	of.	岁	P.	1	À	Å	A	성	ł	9	Comp. Ex.	Comp. Ex	Como Ex	Comp. Ex.								
-	7 2 2 2 3 4 5 7 5 7 5 7 5 7 5 7 5 7 5 7 5 7 5 7 5	Weldability	0	0			0	0	0			0		0			×			0								0
	Plating	adhesion	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	×	0	×	0	V	×	×	×	×	0
	Appearance	of coating	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	×I	0	×	0	⊽	×	×	×	×	×
rapte po	Al content of	coating, %	0.22	0.18	0.28	0.16	0.18	0.82	0.13	0.16	0.25	0.17	0.23	0.32	0.18	0.20	0.21	0.16	0.32	0.13	0.19	0.23	0.28	0.32	0.28	0.32	0.29	0.33
TP T	nt of	coating, %	10	10	1	11	1	10	10	12	1	10	10	11	•		10	10	9	10	10	6	10	•	ı	6	1	11
	tent of	æ	89	89	86	88	94	68	89	87	66	89	68	88	66	93	89	68	93	89	88	90	89	66	94	90	96	88
	Retained	۲٬۰ ۵۶	8	10	13	11	6	6	15	æ	10	7	118	4	15	1	22		8	1	9	-	7	9	2	4	S	2
	E1, &	- 1	36	34	34	31	30	36	31	36	35	36	31	35	32	28	25	28	30	29	33	20	34	33	33	33	33	32
	TS, MPa		630	099	720	640	780	580	840	640	630	620	810	019	830	540	810	570	710	550	620	260	700	640	099	650	999	029
	Sample		-	7	9	4	2	9	7	8	6	ព	=	12	13	14	15	16	17	81	19	20	21	22	23	24	25	26

Note) In the table, mark ____ indicates that the value is outside the scope of the present invention.

Table B6 (continuation from Table B5)

Remarks	Comp. Ex	Comp. Ex.	Comp. Ex	Comp. Ex.																					
Weldability	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Plating	0	0	0	0	0	0	Ø	0	0	⊲	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Appearance of	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Al content of		0.21	0.23	0.63	0.27	0.34	0.28	0.41	0.23	0.23	0.23	0.32	0.23	0.32	1.23	0.41	0.21	0.23	0.63	0.27	0.34	0.41	0.23	0.32	1.23
Fe content of coating %	. 1	10	6	10	10	11	15	10	5	15	7	15	10	10	10	10	10	Ġ	10	10	11	10	10	10	10
Zn content of coating, %	6	89	06	89	89	88	84	88	94	84	92	84	88	89	88	89	89	06	89	68	88	88	89	89	88
Retained Y, %	1	1	1	1	-1	÷1	9	П	5		9		1		7				-	-	-	П	(1	7
E1, %	24	26	20	22	26	24	34	29	35	29	35	28	26	23	35	24	26	20	22	26	24	29	26	23	35
TS, MPa	550	909	620	580	550	650	610	009	610	570	630	580	580	560	630	550	009	620	280	550	650	900	280	260	630
Sample	27	28	29	æ	31	32	33	34	35	36	37	38	39	9	4	42	43	44	45	46	4.1	48	49	22	51

Note) In the table, mark ____ indicates that the value is outside the scope of the present invention.

Example B2

5

10

15

20

25

30

35

Steels having chemical compositions shown in Table B7 and Table B8 (continuation from Table B7) were reheated to 1250°C, were then finish rolled at 900°C, and were coiled at 650°C to prepare 4 mm-thick hot rolled steel sheets. Scale deposited on the surface of the hot rolled steel sheets was removed with hydrochloric acid, and the steel sheets were then cold rolled to prepare 1.4 mm-thick cold rolled steel The cold rolled steel sheets were annealed under conditions shown in Table B9 and Table B10 (continuation from Table B9), were coated, and were then temper rolled with a reduction of 0.5%. The steel sheets thus obtained were subjected to the following performance evaluation tests, that is, "tensile test," "retained austenite measuring test," "welding test," "appearance of coating," "plating adhesion," and "measurement of concentration in coated layer." In the coating in the above production of the steel sheets, both sides of the cold rolled steel sheets were coated at a coverage of coating of 50 g/m^2 per side.

In the "tensile test," a JIS No. 5 tensile test piece was extracted in C-direction, and a cold tensile test was carried out under conditions of gauge thickness 50 mm and tensile speed 10 mm/min.

The "retained austenite measuring test" was carried out by a method called "5-peak" method. In this method, a quarter of the sheet thickness from the surface toward the inner side of the sheet was chemically polished, α -iron intensity and γ -iron intensity were then measured by X-ray diffractometry using an Mo bulb, and the volume fraction of retained austenite was determined based on the α -iron intensity and the γ -iron intensity.

The "welding test" was carried out by performing spot welding under welding conditions of welding current: 10 kA, applied pressure: 220 kg, welding time: 12 cycles, electrode diameter: 6 mm, electrode shape: domed, and tip: 6ϕ -40R, and counting the number of continuous spots provided until the

welding reached the point at which the nugget diameter became below 4√t wherein t represents sheet thickness. The counted number of continuous spots was evaluated according to the following criteria. ⊚: more than 2,000 continuous spots,
5 ⊙: more than 1,000 continuous spots, △: 500 to 1,000 continuous spots, and ×: less than 500 continuous spots. Here ⊚ and ⊙ were regarded as acceptable, and △ and × as unacceptable.

The "appearance of coating" was determined by visually inspecting the appearance of the coated steel sheet for noncoated sites and evaluating the results according to the following criteria.

 \bigcirc : not more than $5/dm^2$, \triangle : 6 to $15/dm^2$, and \times : not less than $16/dm^2$.

15 Here \bigcirc was regarded as acceptable, and \triangle and \times as unacceptable.

The "plating adhesion" was determined by subjecting the coated steel sheet to a 60-degree V bending test, then performing a tape test, and evaluating the results according to the following criteria.

Blackening in tape test (%)

©: 0 to 10

10

20

30

35

O: 10 to less than 20

 \triangle : 20 to less than 30

25 X: not less than 30

(\bigcirc and \bigcirc : acceptable, \triangle and \times : unacceptable)

The "measurement of concentration in coated layer" was carried out by dissolving the coated layer in 5% hydrochloric acid containing an amine-based inhibitor and then analyzing the solution by ICP emission spectroscopy.

The results of the above performance evaluation tests are shown in Table B11 and Table B12 (continuation from Table B11). For all of samples 52 to 64 which are examples of the present invention, the total elongation was not less than 30% while ensuring a tensile strength of not less than 550 MPa, indicating that high strength and good press formability

5

10

15

20

25

30

35

could be simultaneously realized. At the same time, for these samples, the plating adhesion was acceptable.

By contrast, samples 65 to 77, which are comparative examples, could not attain the object of the present invention due to poor strength-ductility balance or poor plating adhesion, because, for sample 65, the content of carbon (C) was lower than the carbon (C) content range specified in the present invention; for sample 66, the content of carbon (C) was higher than the carbon (C) content range specified in the present invention; for sample 67, the content of silicon (Si) was lower than the silicon (Si) content range specified in the present invention; for sample 68, the content of silicon (Si) was higher than the silicon (Si) content range specified in the present invention; samples 69 and 70 failed to satisfy the relationship between silicon (Si) and aluminum (Al) specified in the present invention; for sample 71, the content of manganese (Mn) was lower than the manganese (Mn) content range specified in the present invention; for sample 72, the content of manganese (Mn) was higher than the manganese (Mn) content range specified in the present invention; for sample 73, the content of aluminum (Al) was higher than the aluminum (Al) content range specified in the present invention; for samples 74, 75, and 76, the content of beryllium (Be) + calcium (Ca) + magnesium (Mg) + zirconium (Zr) was lower than the beryllium (Be) + calcium (Ca) + magnesium (Mg) + zirconium (Zr) content range specified in the present invention; and, for sample 77, the content of beryllium (Be) + calcium (Ca)+ magnesium (Mg) + zirconium (Zr) was higher than the beryllium (Be) + calcium (Ca) + magnesium (Mg) + zirconium (Zr) content range specified in the present invention.

For samples 78 to 102 as comparative examples, although the chemical compositions of the steels fell within the scope of the present invention, any one of the treatment conditions was outside the scope of the present invention. These comparative samples suffered from poor strength-ductility balance or poor plating adhesion and thus could not attain the object of the present invention.

Table B7

							Chem	ical co	mositi	Chemical composition, weights	hts			2	Domarke
Steel	U	Si	Mn	ď	s	Z	Be	ß	Mg	Zr	Be+Ca+Mg+Zr	Others	S1+0.8A1		lar. No
aa	0.08	1.21	1.55	0.004	0.005	0.25	0.056	0	0	0	0.056		1.41	Ex. of	of inv.
न्न	0.12	0.24	2.21	0.014	0.003	0.73	0	0.050	0	0	0.050	Mo: 0.11, Bi: 0.02	0.824	Ex. of	of inv.
ac	0.16	1.30	1.40	0.008	0.004	0.02	0	0	0.022	0.022	0.044	Sn: 0.05, Cr: 0.22	1.316	Ex. of	of inv
ad	0.13	0.65	1.00	0.009	0.006	0.74	0	0	0	0.120	0.120	Ni: 0.12	1.242	Ex.	of inv
ae	0.04	1.30	2.40	0.015	0.002	0.21	0.082	0.060	0.043	0.020	0.205	Ce: 0.002, Nb: 0.03	1.468	EX.	of inv.
af.	0.07	0.34	0.90	0.012	0.012	0.65	0.008	0.002	0.008	0.001	0.019	T1: 0.02, Zr: 0.05	0.86	Ex.	of inv.
pre	0.17	1.70	1.50	0.005	0.008	0.23	0.112	0.200	0.100	0.100	0.512	SD: 0.003	1.884	Ex.	of inv
됞	0.09	0.82	1.40	0.005	0.004	0.34	0.187	0	0.742	0.030	0.959	-	1.092	EX.	of inv
교	0.11	0.46	1.60	0.012	0.011	0.95	0.742	0.110	0.046	0.020	0.918	Y: 0.07, Se: 0.01	1.22	Ex.	of inv
a.	0.07	1.12	1.30	0.004	0.005	0.02	0.006	0.006 0.320	0.100	0	0.426	-	1.136	EX.	of inv.
শ্ব	0.18	0.93	1	1.6 0.008	0.009	0.14	0	0.230	0.230	090.0	0.520	Mo: 0.04, Ti: 0.01, Sb: 0.02	1.042	EX.	Ex. of inv.
급	0.08	0.82	1.70	0.004	0.005	0.13	0.046	0.046 0.020	0	0.010	0.076	•	0.924	Ex.	of inv.
Ħ	0.17	1.40	1.70	1.70 0.005	0.008	0.23	0	0.200	0.102	0.100	0.402	Co: 0.05	1.584	Ex.	of inv.

In the table, mark ____indicates that the value is outside the scope of the present invention; and mark * indicates that the value does not satisfy the relationship between Al and Si specified in the present invention; and mark -- indicates that the constituent is not contained. Note)

Table B8 (continuation from Table B7)

	Remarks	The state of the s	COMP. EA.	Comp. Ex.	Comp. Ex.	Comp	Comp Ex	Comp. E.A.	Comp. Ex.	Comp. Ex.	Comp Dv		. w	Comp. Ex.	Comp Pir		Comp. EX.	Comp. Ex.	
	S1+0 8A1	a. 0	0,.0	0. /96	0.442	2.112	0 316	25.6	7.134	0.536	1 162	1 844	F. C. T.	0.876	0.876	910	0.0.0	0.876	
	Others	1		sn: 0.05	Ce: 0.2		V: 0.01 2r: 0.02			Cu: 0.22, B1: 0.021	Cr: 0 23 Mo: 0 09			Ti: 0.03, Nb: 0.03	Ni: 0.11. Sb: 0.02	80.00	20:0		that the value is outside the scope of the present importion.
11%	Be+Ca+Ma+Zr	0.088	730 0	0.234	0.073	0.034	0.572	0 402	2010	0.332	0.360	0.051		0.001	0.003	0 002		1.402	ne of the nre
Chemical composition, weights	12	0.030	000	0.020	0	0.002	0.230			0.220	0	0.023		5	0	0.001		0.449	the sco
mpositic	Mg	0.020	0 112	7:176	0.010	0.030	0.112	c	•	0.008	0.130	0.010	í	0	0.001	0		7:17	outside
mical co	Ca	0.010	0000	0.020	0.020	0.002	0.020	0.080		0.042	0.230	0.010	,	>	0.002	0	000	0.2.0	alue is
Che	Be	0.028	0 102	- 1	0.043	0	0.210	0.192		0.062	0	0.008	000	- 1	0	0.001	┸	7.00.7	at the v
	Z	0.55	0 22		0.39	0.24	*0.12	*0.73		77.0	0.54	1.63	33	75.0	0.32	0.32	233	25.0	
	S	0.005	0 002		- 1	0.002	0.007	0.003	100	0.003	0.005	0.006	000	- 1	0.002	0.002	200	- 1	indicates
	P	0.003	0.013		0.00/	0.008	0.004	0.005	000	0.008	0.009	0.005	1 22 0 012		0.012	0.012	0 012		¥
	된	1.03	1.82		1.34	0.97	0.58	1.52	9,0	3	2.65	0.85	1 22		77.1	1.22	1 22		e, ma
	SI	0.34	0.62		1	- 1	*0.22	*1.55	75.0	1	0.73	0.54	0 62	- 1.	0.07	0.62	0.62		In the table, mark
	ပ	0.01	0.22	;	?	0.16	0.15	0.12	90	3	0.14	0.12	60.0	3	5	60.0	0.09		
Steel		an	ao	!	}	ad	교	9.8	÷		ag g	å	36		ş	ay	az		Note)

In the table, mark ____indicates that the value is outside the scope of the present invention; mark * indicates that the value does not satisfy the relationship between Al and Si specified in the present invention; and mark -- indicates that the constituent is not contained.

	_			-	-		_	_	1	_	_	_		-	_	-	,	,	_	1	_	_	·	-	$\overline{}$	т-	-1	_
		Remarks	8	4	15	5	10	5	3	8	19	3	8	8	ij	19	8	8	8	8	8	প্ত	8	15	5	8	1	5 6
	Cooling	temp.,	7 68 1	150	180	180	180	180	180	180	180	180	180	180	200	180	180	180	180	180	180	180	180	180	180	180	180	1 BO
	Cooling	rate,	10	15	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	10	5
	Alloying	time,	25	30		25	-	,	30	25	•	25	30	10			15	25	30	20	25	25	25		1	25		25
	Alloying	temp.,	500	520		500			200	200		200	200	550	,		520	500	580	500	200	500	500	1		200		500
	Souttent 31 content		0.11	0.14	0.16	0.10	0.12	0.15	0.15	0.20	0.11	0.15	0.20	0.18	0.20	0.15	0.14	0.10	60.0	0.16	0.13	0.18	0.17	60.0	0.14	0.14	0.14	0.14
	20.4400	temp., °c	450	450	470	450	440	450	450	450	450	450	450	450	460	450	460	450	450	450	450	450	450	450	440	440	440	440
Table B9	Holding	time,	9		420	,	300		.,	40	200	100	•	9	•	30	-	-	30	1	300	•	30	90	,	•	80	
Tab	Holding	temp. before	400 - 450	١.	360 - 400	,	350 - 380	-	•	400 - 470	360 - 440	400 - 500	1	360 - 440	•	400 - 470	1	-	430 - 500	•	370 - 440	•	400 - 490	400 - 470		•	400 - 470	1
	Cooling	termination	400	400	420	380	370	480	410	440	360	480	440	400	430	410	420	370	480	410	440	420	480	490	400	400	400	440
	Cooling	rate,	10	20	40	130	3	20	20	7	10	20	S	10	15	20	9	20	20	20	6	20	22	20	ъ	10	S	20
	Annealing	time, sec	150	90	100	90	150	90	200	85	150	06	110	120	200	90	120	110	90	90	150	70	90	200	120	110	120	70
	Annealing	temp., °c	800	750	800	750	780	800	750	850	750	800	750	700	900	830	800	750	820	750	870	750	830	750	800	750	800	800
		Steel	88	q	ac	ad	ae	a£	ag	윰	1e	aj	¥	Te	ā	ur	ao	ďe	be	ar.	a.s	at	an	av	WE	¥	ay	2E
		Sample	52	53	54	55	56	57	88	59	09	19	62	63	64	63	99	- 1	89	69	۶	2	72	73	74	27	76	77
-															_	_									_			

Note) In the table, mark ____indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and

GA represents a hot-dip galvannealed steel sheet, and GI represents a hot-dip galvanized steel sheet.

Table B10 (continuation from Table B9)

		Annoy 1 4 n.c.	7,100	Cooling	Cooling	Holding	Holding	Coating	Al content	Alloying	Alloying	Cooling	Cooling	
Sample	Steel	temp	time sec	rate,	termination	temp. before	time,	temp.,	of zinc	temp.,	time,	rate,		Remarks
		, , ,	200 / 2000	°C/sec	temp., °c	coating, °C	sec	ပ္စ	bath, &	ູບ	sec	°C/sec	ပ္စ	
78	aa	600	ک 06	20	360	•	,	470	90.0	500	30	10	180	8
79	aa	950	06	90	480	400 - 500	09	450	0.11	480	50	10	180	8
90	aa	750	S	20	440	430 - 500	20	450	0.20	500	30	10	180	8
81	aa	800	400	20	410	-	•	480	0.17	500	40	20	220	8
82	aa	750	06	н	370	430 - 500	30	450	0.13	510	30	10	180	క
83	ee	900	110	10	300	370 - 440	300	450	0.09	480	50	01	180	8
84	23	770	96	70	520	370 - 440	300	450	0.14	200	30	0,7	180	8
82	aa	830	150	10	420	400 - 490	650	480	0.18	200	30	01	180	8
86	aa	800	70	20	410	400 - 470	80	430	0.11	430	40	15	180	5
87	aa	750	90	25	440	370 - 440	140	480	0.16	620	20	10	100	89
88	aa	850	9	20	420	•	-	450	0.20	500	6	8	180	45
89	ee	750	90	90	480	•	•	450	0.12	200	130	92	180	\$
90	аа	820	70	50	490	400 - 470	250	440	0.10	500	25	m	180	శ్ర
91	aa	750	100	20	360	•	-	450	0.08	200	30	10	300	45
92	aa	830	90	20	480	400 - 500	90	450	0.82	200	25	10	180	f
93	23	909	90	20	360	-	1	470	90.0	-		10	180	5
94	aa	950	06	80	480	400 - 500	09	450	0.11	1	1	10	180	15
95	aa	750	5	20	440	430 - 500	20	450	0.20	ě	-	10	180	15
98	aa	800	420	20	410	1	'	480	0.17	1	-	20	220	15
97	aa	750	06	7	370	430 - 500	30	450	0.13	•	-	10	180	ij
8	aa	800	110	22	300	370 - 440	300	450	0.09		•	10	180	15
66	33	830	150	10	420	400 - 490	720	480	0.18	•	•	10	180	19
100	aa.	820	70	50	490	400 - 470	250	440	0.10	,		б	180	5
101	aa	750	100	20	360	-	•	450	0.08	•		10	300	5
102			06	20	480	400 - 500	9	450	0.82			10	180	15
Note)	In th	the table, n	mark in	indicates	that the va	that the value is outside	the	scope of	the present	t invention;	on;			

In the table, mark indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and GA represents a hot-dip galvannealed steel sheet, and GI represents a hot-dip galvanized steel sheet.

Table B11

<u>"</u>	5.	inv.	ņ.	inv.	inv.													Ī								
Remarks	of inv.	å	of	of	• of	of	of	of	of	of	of inv	of	of	Comp. Ex.												
~	EX	EX.	EX	Ä	EX.	EX	Ex.	EX	EX.	ΕX	Ex.	EX.	Ex.	ပိ	S	S	ပ္ပ	ပ္ပိ	ပိ	S	ည်	ខ្ល	ပ္ပိ	ပ္ပိ	ပိ	
Weldability	0	0	0	0	0	0	0	0	0	0	0	0	0	0	×I	0	0	0	0	0	×	0	0	0	0	
Plating adhesion	0	0	0	0	0	0	0	0	©	0	0	0	0	0	0	0	×I	0	×	0	⊽	×I	×I	×I	×	
Appearance of coating	0	0	0	Ó	0	0	0	0	0	0	0	0	0	0	0	0	×	0	×I	0	ಶ	×I	×I	×	×I	
Al content of coating, %	0.22	0.18	0.28	0.16	0.18	0.82	0.13	0.16	0.25	0.17	0.23	0.32	0.18	0.20	0.21	0.16	0.32	0.13	0.19	0.23	0.28	0.32	0.28	0.32	0.29	
Fe content of coating, %	10	10	-	11	•	10	10	12	_	10	. 01	11	-	_	10	10	9	10	10	6	10	-	•	6	1	
Zn content of coating, %	89	68	86	88	76	68	68	87	66	68	89	88	66	93	89	89	93	89	89	06	89	66	94	06	96	
Retained 7, 8	8	10	13	11	3	6	15	8	10	7	18	Þ	15	1	22	-	8	-1	9		7	9	2	4	5	
El, &	36	34	34	31	30	36	31	36	35	36	31	35	32	28	25	28	30	29	33	20	34	33	33	33	33	
TS, MPa	630	099	720	640	780	580	840	640	630	620	810	610	830	540	810	570	710	550	620	560	700	640	099	650	665	
Sample	52	53	54	55	56	57	58	59	09	61	62	63	64	65	99	67	68	69	70	71	72	73	74	75	76	

indicates that the value is outside the scope of the present invention. Note) In the table, mark

Table B12 (continuation from Table B11)

			1		ži į	×.	EX.	X.	Ä.	Ex.	EX.	EX.	Ex.	EX.	2	2		3 6	ž	×	EX.	X.	Ex.	Ex.	E.	X	×	, and	Ī
		Remarks	100		Comp	Xa diloo	Comp. Ex.	Comp. Ex	Comp					Comp. Ex	Como. Ex	Comp. Ex	Comp. Ex												
		Weldability	@	0	9 @	9	9	0	0	0	0	0	0	0	С	@	0	C				0	0	0	0	0	0	0	
	Plating	adhesion	@	0	0		9	9	9	7	0	0	⊲!	0	۵	0	0	0	0	0	9 @	D	0	0	0	0	0	0	
(דום ביו	Appearance of	coating	C							0	0	0	0	0	0	0	0	0	C	C	þ		2	Ó	0	0	0	0	C
יייי ביייי ביייי דמייי דמייי דמייי	Al content of	coating, &	0.41	0.21	0.23	69 0	20.0	77.0	40.0	0.20	0.41	0.23	0.23	0.23	0.32	0.23	0.32	1.23	0.41	0.21	0 23		0.03	0.27	0.34	0.41	0.23	0.32	
	Fe content of	coating, &	10	10	6	10	10	-	21 21	2 5	07	20	15	7	15	10	10	10	10	10	6	9.	2	10	11	10	10	10	
	Zn content of	coating, %	89	89	06	89	89	88	84	68	S	94	84	92	84	89	89	88	68	89	06	8	3	80	90	88	68	68	88
	Retained	۲, 8	1	1	T	1	н		9	-	(I U	,	,	۰	-		-	7	н	٦	1	П	-	-	1.	40,	-	- 1	_
	E1, 8		24	26	20	22	26	24	34	29	35	300	2/2	20	87	26	23	35	24	26	20	22	26	24	,	67	27	52	ς,
	TS, MPa		550	009	620	580	550	650	610	909	610	570	063	000	280	280	260	630	550	909	620	580	550	650	009	200	200	200	000
	Sample		78	79	80	18	82	83	84	85	86	87	ä	3 8	6	3	16	25	93	94	95	96	97	86	8	15		195	707

Note) In the table, mark ____ indicates that the value is outside the scope of the present invention.

Example B3

5

10

15

20

25

30

35

Steels having chemical compositions shown in Table B13 and Table B14 (continuation from Table B13) were reheated to 1250°C, were then finish rolled at 900°C, and were coiled at 650°C to prepare 4 mm-thick hot rolled steel sheets. Scale deposited on the surface of the hot rolled steel sheets was removed with hydrochloric acid, and the steel sheets were then cold rolled to prepare 1.4 mm-thick cold rolled steel The cold rolled steel sheets were annealed under conditions shown in Table B15 and Table B16 (continuation from Table B15), were coated, and were then temper rolled with a reduction of 0.5%. The steel sheets thus obtained were subjected to the following performance evaluation tests, that is, "tensile test," "retained austenite measuring test," "welding test," "appearance of coating," "plating adhesion," and "measurement of concentration in coated layer." In the coating in the above production of the steel sheets, both sides of the cold rolled steel sheets were coated at a coverage of coating of 50 g/m^2 per side.

In the "tensile test," a JIS No. 5 tensile test piece was extracted in C-direction, and a cold tensile test was carried out under conditions of gauge thickness 50 mm and tensile speed 10 mm/min.

The "retained austenite measuring test" was carried out by a method called "5-peak" method. In this method, a quarter of the sheet thickness from the surface toward the inner side of the sheet was chemically polished, α -iron intensity and γ -iron intensity were then measured by X-ray diffractometry using an Mo bulb, and the volume fraction of retained austenite was determined based on the α -iron intensity and the γ -iron intensity.

The "welding test" was carried out by performing spot welding under welding conditions of welding current: 10 kA, applied pressure: 220 kg, welding time: 12 cycles, electrode diameter: 6 mm, electrode shape: domed, and tip: 6ϕ -40R, and counting the number of continuous spots provided until the

welding reached the point at which the nugget diameter became below $4\sqrt{t}$ wherein trepresents sheet thickness. The counted number of continuous spots was evaluated according to the following criteria. \odot : more than 2,000 continuous spots, \odot : more than 1,000 continuous spots, Δ : 500 to 1,000 continuous spots, and \times : less than 500 continuous spots. Here \odot and \odot were regarded as acceptable, and Δ and \times as unacceptable.

The "appearance of coating" was determined by visually inspecting the appearance of the coated steel sheet for noncoated sites and evaluating the results according to the following criteria.

 \bigcirc : not more than $5/dm^2$, \triangle : 6 to $15/dm^2$, and \times : not less than $16/dm^2$.

15 Here \bigcirc was regarded as acceptable, and \triangle and \times as unacceptable.

The "plating adhesion" was determined by subjecting the plated steel sheet to a 60-degree V bending test, then performing a tape test, and evaluating the results according to the following criteria.

Blackening in tape test (%)

©: 0 to 10

10

20

25

30

35

O: 10 to less than 20

 \triangle : 20 to less than 30

X: not less than 30

(\bigcirc and \bigcirc : acceptable, \triangle and \times : unacceptable)

The "measurement of concentration in coated layer" was carried out by dissolving the coating layer in 5% hydrochloric acid containing an amine-based inhibitor and then analyzing the solution by ICP emission spectroscopy.

The results of the above performance evaluation tests are shown in Table B17 and Table B18 (continuation from Table B17). For all of samples 103 to 115 which are examples of the present invention, the total elongation was not less than 30% while ensuring a tensile strength of not less than 550 MPa, indicating that high strength and good press formability

5

10

15

20

25

30

35

could be simultaneously realized. At the same time, for these samples, the plating adhesion was acceptable.

By contrast, samples 116 to 128, which are comparative examples, could not attain the object of the present invention due to poor strength-ductility balance or poor plating adhesion, because, for sample 116, the content of carbon (C) was lower than the carbon (C) content range specified in the present invention; for sample 117, the content of carbon (C) was higher than the carbon (C) content range specified in the present invention; for sample 118, the content of silicon (Si) was lower than the silicon (Si) content range specified in the present invention; for sample 119, the content of silicon (Si) was higher than the silicon (Si) content range specified in the present invention; samples 120 and 121 failed to satisfy the relationship between silicon (Si) and aluminum (Al) specified in the present invention; for sample 122, the content of manganese (Mn) was lower than the manganese (Mn) content range specified in the present invention; for sample 123, the content of manganese (Mn) was higher than the manganese (Mn) content range specified in the present invention; for sample 124, the content of aluminum (Al) was higher than the aluminum (Al) content range specified in the present invention; for samples 125, 126, and 127, the content of scandium (Sc) + yttrium (Y) + lanthanum (La) + cerium (Ce) was lower than the scandium (Sc) + yttrium (Y) + lanthanum (La) + cerium (Ce) content range specified in the present invention; and, for sample 128, the content of scandium + yttrium + lanthanum + cerium was higher than the scandium + yttrium + lanthanum + cerium content range specified in the present invention.

For samples 129 to 153 (see Table B18) as comparative examples, although the chemical compositions of the steels fell within the scope of the present invention, any one of the treatment conditions was outside the scope of the present invention. These comparative samples suffered from poor strength-ductility balance or poor plating adhesion and thus

could not attain the object of the present invention.

Table B13

										_	_	_	_	_	_		_		_		.	
	Remarks		Ex. of inv.	of inv.	The Of Start	01 THY:	or inv.	Ex. of inv.	ישני שלי	;	or Inv.	of inv.		Ex. of Inv.	of inv		Ex. of inv.		Ex. of inv.	9	EX. OL TIN.	
	_		ద	KX	i e	3	Ë	Ä	Ġ	1	EX.	EX.	i	<u> </u>	à	ij	<u> </u>		E	ţ	3	
	500	S1+0.8AL	1.41	0 R24	1 346	1.310	1.242	1.468	90 0	00.0	1.884	1 092	3	1.22	1 126	25.7	1.042		0.924		1.364	
!		Others	•	20 0 11 0 00	MO: 0.11, Ca. 0.02	sn: 0.05, Cr: 0.22	Ni: 0.12	Sb: 0.002, Nb: 0.03	20 0	T1: 0.02, &F: 0.03	Mg: 0.003			Bi: 0.07, Ca: 0.01			Mo: 0.04, Ti: 0.01,	Mg: 0.02			Mg: 0.02, Co: 0.05	
4+4	211	Sc+X+La+Ce	0.056		0.030	0.044	0.120	0 205		0.019	0.512	020	0.939	0 918		0.426	000	0.350	200	0.0.0	0.402	
2,000	JII, WELY	çe	0	,	٥	0.022	0.120	0 000	0:020	0.001	0.100		0.030	0 00	20.0	0	0	0.0	3	0.010	0.100	
Chemical composition, weights	TOTSOGIE	Ľa	c	,	0	0.022	0	0 043	2	0.008	0.100		0.742	0 046	2.0.0	0.100	0.0	0.230	,	0	0.102	
	Ical co	X	6	0.050			0	ļ	200.0	0.008 0.002	0000		0	0110	777.	0.320		0.230		0.046 0.020	0.200	
	CDE	Sc	9			0	6	000	0.002	0.008	0 112	7	0.187	242	0.144	0.006		0		0.046	0	,
		Z	S			0.02	14		0.21	0.65	0 23	ı	0.34	6	0.95	0.02	ı	0.14	1	0.13	0 23	
		S	03			0.004	1	- 1	0.002	0.012	1		0.004		110.0	0.005	1.6 0.008 0.009			0.005	000	
		4		0.004	0.014	0.008	1	٠.	0.015	0.012	ı	0.003	0 005		0.012	0.004		0.008		1.70 0.004	1 70 0 005	} > >
		¥	1	1.55	2.21	1 40	1	- 1	2.40	06 0	•	1.30	1 40	1	1.60	1.30	1		-		1 70	>
		10	1	1.21	0.24	1 30	1	- 1	1.30	DF 0	1	٥/٠٢	0 82	- 1	0.46	1 12	L	0.93		0.82	1	
	_		,	0.08	0.12	0 16		0.13	0.04	0 07		0.17	00		0.11	0.07		0.18		0.08	1.	-
		Steel		ď	2	٢	3 3	8	þe	4	1	ğ	1		ቯ	ž		ጟ		7		Ħ

Table B14 (continuation from Table B13)

							Cher	nical co	mpositic	Chemical composition, weight%	nt8,			Remarks
Si Mn P S Al	Si Mn P S Al	P S Al	s Al	¥	_		Sc	Y	La	S.	Sc+Y+La+Ce	Others	Si+0.8Al	
0.01 0.34 1.03 0.003 0.005 0.55 0	0.34 1.03 0.003 0.005 0.55	0.003 0.005 0.55	0.003 0.005 0.55	0.005 0.55	0.55		0.028	0.010	0.020	0.030	0.088	1	0.78	Comp. Ex.
0.22 0.62 1.82 0.013 0.002 0.22 0	1.82 0.013 0.002 0.22	0.013 0.002 0.22	0.013 0.002 0.22	0.002 0.22	22	0	0.102	0.020	0.112	0.020	0.254	Sn: 0.05	0.796	Comp. Ex.
0.13 0.13 1.34 0.007 0.003 0.39 0.	0.007 0.003 0.39	0.007 0.003 0.39	0.007 0.003 0.39	0.003 0.39	39	0.	0.043	0.020	0.010	0	0.073	Se: 0.2	0.442	Comp. Ex.
0.16 1.92 0.97 0.008 0.002 0.24	1.92 0.97 0.008 0.002 0.2	0.008 0.002 0.2	0.008 0.002 0.2	0.002 0.2	~		0	0.002	0.030	0.002	0.034	•	2.112	Comp. Ex.
0.15 *0.22 0.58 0.004 0.007 *0.12 0.	*0.22 0.58 0.004 0.007 *0.12	0.58 0.004 0.007 *0.12	0.004 0.007 *0.12	0.007 *0.12	. 12		0.210	0.020	0.112	0.230	0.572	V: 0.01, Zr: 0.02	0.316	Comp. Ex.
*1.55 1.52 0.005 0.003 *0.	*1.55 1.52 0.005 0.003 *0.73	1.52 0.005 0.003 *0.73	0.005 0.003 ±0.73	0.003 +0.73	.73		192	0.080	0	0.130	0.402		2.134	Comp. Ex.
0.06 0.36 0.18 0.008 0.003 0.22 0.062	0.36 0.18 0.008 0.003 0.22	0.18 0.008 0.003 0.22	0.008 0.003 0.22	0.003 0.22	22	0.0	12	0.042	0.008	0.220	0.332	Cu: 0.22, Ca: 0.021	0.536	Comp. Ex.
0.14 0.73 2.65 0.009 0.005 0.54 0	0.73 2.65 0.009 0.005 0.54	0.009 0.005 0.54	0.009 0.005 0.54	0.005 0.54	0.54	0		0.230	0.130	0	0.360	Cr: 0.23, Mo: 0.09	1.162	Comp. Ex.
0.12 0.54 0.85 0.005 0.006 1.63 0.008	0.54 0.85 0.005 0.006 1.63	0.005 0.006 1.63	0.005 0.006 1.63	0.006 1.63		0.0	80	0.010	0.010	0.023	0.051	•	1.844	Comp. Ex.
0.09 0.62 1.22 0.012 0.002 0.32 0.001	0.62 1.22 0.012 0.002 0.32	1.22 0.012 0.002 0.32	0.012 0.002 0.32	0.002 0.32	32	0.0	딩	0	0	0	0.001	Ti: 0.03, Nb: 0.03	0.876	Comp. Ex.
0.09 0.62 1.22 0.012 0.002 0.32 0	0.62 1.22 0.012 0.002 0.32	1.22 0.012 0.002 0.32	0.012 0.002 0.32	0.002 0.32	.32		0	0.002	0.001	0	0.003	Ni: 0.11, Mg: 0.02	0.876	Comp. Ex.
0.09 0.62 1.22 0.012 0.002 0.32 0.001	0.62 1.22 0.012 0.002 0.32	0.012 0.002 0.32	0.012 0.002 0.32	0.002 0.32	32	0	70	0	0	0.001	0.002	Sn: 0.04	0.876	Comp. Ex.
0.09 0.62 1.22 0.012 0.002 0.32 0.8	0.62 1.22 0.012 0.002 0.32	0.012 0.002 0.	0.012 0.002 0.32	0.002 0.32	32	0.8	19	0.200	0.112	0.229	1.402		0.876	Comp. Ex.

		-	_	_	1	т-	_	_	i			_		_							-				_				_	
		Remarks		f	3	ថ	8	10	15	3	3	5	8	8	5	5 6	15	3	5 6	\$ 8	3	5 8	5 6	5 6	3	GI.	ម	8	15	8
	Cooling	temp.,	ູບ	180	150	180	180	180	180	180	180	180	180	180	180	200	180	180	180	180	180	180	180	1 5		8	180	180	180	180
	Cooling	rate,	°C/sec	10	15	10	01	10	10	2	10	10	2	2	2	S	20	2	07	10	97	2	2	5		20	10	10	10	51
	Alloying	time,	sec	25	30	•	25	-	•	30	25	1	25	30	ន្ព			15	25	e	20	25	25	25		1	•	25	ı	25
	ng	·	1	200	520	•	200	-	'	200	500	•	200	500	550	•	•	520	500	580	500	500	500	200		1		200	•	200
	Al content of	od zinc bath, &		0.11	0.14	0.16	0.10	0.12	0.15	0.15	0.20	0.11	0.15	0.20	0.18	0.20	0.15	0.14	0.10	0.09	0.16	0.13	0.18	0.17	0.09		0.14	0.14	0.14	0.14
<u> </u>	Coating	temp., °d 2		420	450	470	450	440	450	450	450	420	450	450	450	460	430	460	450	450	450	450	450	450	450	440		0840	440	440
Table B15	Holding time		3	2	•	420	•	300	•	.	Q	200	100	•	09	•	8	•	•	8	•	900	'	8	80		1	1 8	20	.
	Holding temp. before	coating. °C	400 - 450	· [·	• [300 - 400	•	350 - 380	-	•	360 - 470	1	400 - 200	۱ ا	300 - 440	•	400 - 470	:	1	430 - 500	•	3/0 - 440	•	•	400 - 470	•	•	400 - 470	٠ ا	"
27,100	rion -		400	400	430	280	230	370	410	440	360	400	940	400	730	430	007	270	0/5	480	240	250	075	084	490	400	400	400	440	that the walne
Cool that		"C/sec	_	20	6	150	-	200	202	7	S	20	6	, 2	=	200	2	, =	2 5	200	9	200	2 5	2 8	23	5	10	80	202	89
	Annealing	רייוופי פכנ	150	06	100	06	150	90	200	85	150	06	110	120	200	06	120	110	06	90	150	70	6	200		120	110	120	70	mark inc
	Annealing temp. °c	,	800	. 750	800	750	780	800	750	850	750	800	750	700	800	830	800	750	820	750	870	750	830	750	000	900	750	800	800	table,
	Steel	T	ğ	qq	pc	Þq	Bg Z	ρţ	Бq	ųq	P1	દિવ	γq	Ιq	Eq	pu	ନ୍ଦ	ďq	ď	br	bs	Þţ	Pa Pa	ž	3	†	ŭ	à	zq	In the
	Sample		103	104	105	106	107	108	109	110	111	112	113	114	115	116	117	118	119	120	121	122	123	124	125		126	127	128	Note)

GA represents a hot-dip galvannealed steel sheet, and GI represents a hot-dip galvanized steel sheet. In the table, mark indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and

ð

Table B16 (continuation from Table B15)

									/200					
•		Annealing	Annealing	Cooling	Cooling	Holding	Holding	Coating	Al content	Alloying	Alloying	Cooling	Cooling	
Sample	Steel	temo. °C	time sec	rate,	termination	temp. before	time,	temp.,	of zinc	temp.,	time,	rate,	temp.,	Remarks
			200	°C/sec	temp., °c	coating, °C	sec	္ရင	bath, 8	္စ	580	°C/sec	ပ္စ	
129	ba	600	90	20	360	-	•	470	0.08	200	30	10	180	5
130	ba	950	96	90	480	400 - 500	60	450	0.11	480	50	97	180	3
131	ba	750	មា	20	440	430 - 500	20	450	0.20	200	30	01	180	\$
132	ba	800	400	20	410	1	•	480	0.17	500	40	20	220	\$
133	ba	750	90		370	430 - 500	30	450	0.13	510	30	10	180	5
134	Ъа	800	110	10	300	370 - 440	300	450	60.0	480	30	0.1	180	85
135	ba	770	90	70	520	370 - 440	300	450	0.14	500	30	10	180	\$
136	ba	830	150	10	420	400 - 490	630	480	0.18	500	30	2	180	89
137	ba	800	70	20	410	400 - 470	08	430	0.11	430	40	15	180	45
138	ba	750	90	25	440	370 - 440	140	480	0.16	620	20	10	100	G.
139	pa	850	9	20	420	-	-	450	0.20	500	6		180	3
140	ba	750	06	80	480	1	•	450	0.12	500	130	10	180	8
141	ba	820	70	50	490	400 - 470	250	440	0.10	500	25	3	180	8
142	ba	750	100	20	360	_	-	450	80.0	500	30	93	300	5
143	pa	830	06	20	480	400 - 500	90	430	0.82	500	25	10	180	8
144	ba	909	06	20	360	•		470	90.0		1	9	180	i
145	ра	950	06	90	480	400 - 500	09	450	0.11	,	,	10	180	15
146	þa	750	ξ.	20	440	430 - 500	20	450	0.20	,		10	180	15
147	pg	800	420	20	410	1	,	480	0.17	•		20	220	15
148	pa	750	06	1	370	430 - 500	30	450	0.13	1		10	180	5
149	ba	800	110	10	300	370 - 440	300	450	0.09	-	,	10	180	15
150	pa	830	150	10	420	400 - 490	720	480	0.18	•		ខ្ម	180	GI
151	pa	820	70	50	490	400 - 470	250	440	0.10	1		9	180	ij
152	pa	750	100	20	360	1	•	450	90.0	1	'	2	300	GI
153	pa	830	90	20	480	400 - 500	09	450	0.82	,		10	180	IS
Note)	In t	the table.	mark in	indicates	that the wa	traine is outestable	44 06	30 000	1					

In the table, mark indicates that the value is outside the scope of the present invention; the heating rate after coating was constant and 10°C/sec; and GA represents a hot-dip galvannealed steel sheet.

Table B17

Retained In content 7, 8 coating, 8	 	ent of	Fe content of coating, %	of Al content of coating, %	Appearance of coating	Plating adhesion	Weldability	Remarks
-	89		10	0.22	0	0	0	Ex. of inv.
-	89		10	0.18	0	0	0	Ex. of inv.
34 13 98	86		-	0.28	0	0	0	Ex. of inv.
31 11 88	88	ļ	11	0.16	0	0	0	Ex. of inv.
30 3 94	94		1	0.18	0	0	0	å
36 9 89	89	-	10	0.82	0	0	0	Ex. of inv.
31 15 89	. 89	I	10	0.13	0	0	0	Ex. of inv.
36 8 87	87	l	12	0.16	0	0	0	Ex. of inv.
35 10 99	66		1	0.25	0	0	0	Ex. of inv.
36 7 89	89	ŀ	10	0.17	0	0	0	of
31 18 89	89		10	0.23	0	0	0	of
35 4 88	88		11	0.32	0	0	0	of
32 15 99	66		•	0.18	0	0	0	Ex. of inv.
-	93		ı	0.20	0	0	0	Comp. Ex.
25 22 89	88	1	10	0.21	0	0	×I	Comp. Ex.
28 1 89	89		10	0.16	0	0	0	Comp. Ex.
30 8 93	93	- 1	9	0.32	×	×I	0	Comp. Ex.
	89	- }	10	0.13	0	0	0	Comp. Ex.
33 6 89	89		10	0.19	×	×I	0	Comp. Ex.
_	06	- 1	6	0.23	0	0	0	Comp. Ex.
34 7 89	.68	-	10	0.28	⊽	۵	×	Comp. Ex.
33 6 99	66		1	0.32	×I	×	0	Comp. Ex.
33 5 94	94		1	0.28	×I	×	0	Comp. Ex.
33 4 90	06	- 1	6	0.32	×I	×I	0	Comp. Ex.
	96	- 1	•	0.29	×I	×I	0	Comp. Ex.
32 5 88	88	- i	11	0.33	×	0	0	Comp. Ex.

Note) In the table, mark ____indicates that the value is outside the scope of the present invention.

Table B18 (continuation from Table B17)

	<u>ي</u> د.	3	X	i i		X	4	1	3 2			<u> </u>					×	2		×	×	×	Š	X	×		T
	Sylemon		Comp. Ex	Comp	Comp. F.v	Como. Ex	Comp.		College	Comp	Comp	Comp	Comp Ex		Coll	Comp. Fr	Comp. Ex	Comp F.v	Como Ex	Comp. Ex	Como. Ex	Comp. Ex	Comp. Ex	Comp. Ex	Comp. Ex	Comp	1
	Weldability	In a second	0	6	0	0	6	0	C	6	C	C	C	C	0	0	0	С	0	0	0	0	0	0	0	С	C
	Plating	adhesion	0	·©	0	0	0	@		0	©	٥	0	<	0	0	0	0	0	0	0	0	0	0	0	0	@
1, 22	Appearance of	coating	0	0	0	0	0	0	0	0	0	0	O	0	0	0	0	0	0	0	0	0	0	.0	0	0	С
	•	coating, %	0.41	0.21	0.23	0.63	0.27	0.34	0.28	0.41	0.23	0.23	0.23	0.32	0.23	0.32	1.23	0.41	0.21	0.23	0.63	0.27	0.34	0.41	0.23	0.32	1.23
	9	coating, *	10	10	6	10	10	11	15	10	ស(1.5	7	15	10	10	10	10	10	6	10	10	11	10	10	10	.01
	ten ,	coating, *	89	89	90	89	89	88	84	89	94	84	92	84	68	89	88	68	. 89	06	89	89	88	89	89	89	88
	Retained	ړ, څ	+1	1	-1	-	1	T	.9	7	S	1	9	-	-1	-1	7	1	1	-1	1			-1	7	-1	7
	E1, &		24	26	20	22	26	24	34	29	35	29	35	28	26	23	35	24	26	20	22	26	24	29	26	23	35
	TS, MPa		550	909	620	280	550	650	610	900	610	570	630	580	580	260	630	550	900	620	280	550	650	000	280	260	630
	Sample		129	130	131	132	133	134	135	136	137	138	139	140	141	142	143	144	145	146	147	891	500	PC :		152	153

Note) In the table, mark ____ indicates that the value is outside the scope of the present invention.

CLAIMS

1. A high strength hot-dip galvanized or galvannealed steel sheet having improved plating adhesion and press formability, said steel sheet comprising:

(a) a steel sheet substrate comprising, by weight,

carbon (C): 0.05 to 0.2%,

silicon (Si): 0.2 to 2.0%,

manganese (Mn): 0.2 to 2.5%, and

aluminum (Al): 0.01 to 1.5%,

said silicon and said aluminum having a mutual relationship represented by formula

 $0.4 (\%) \le Si + 0.8Al (\%) \le 2.0\%,$

said steel sheet substrate further comprising at least one member selected from the group consisting of

- (i) 0.003 to 1.0% of tin (Sn),
- (ii) 0.005 to 1.0% in total of at least one member selected
 from antimony (Sb), bismuth (Bi), and selenium (Se),
- (iii) 0.005 to 1.0% in total of at least one member selected from beryllium (Be), magnesium (Mg), calcium (Ca), and zirconium (Zr), and
- (iv) 0.005 to 1.0% in total of at least one member selected from scandium (Sc), yttrium (Y), lanthanum (La), and cerium (Ce), with the balance consisting of iron (Fe) and unavoidable impurities, the volume fraction of retained austenite in the steel structure being 2 to 20%; and
- (b) a zinc (Zn) coated layer provided on said steel sheet substrate.
- 2. The hot-dip galvanized or galvannealed steel sheet according to claim 1, wherein said steel sheet substrate further comprises, by weight, at least one member selected from nickel (Ni): not more than 2.0%, copper (Cu): not more than 2.0%, and cobalt (Co): less than 0.3%.
- 3. The hot-dip galvanized or galvannealed steel sheet according to claim 1 or 2, wherein said steel sheet substrate further comprises, by weight, at least one member selected

from molybdenum (Mo): less than 0.5%, chromium (Cr): less than 1.0%, vanadium (V): less than 0.3%, titanium (Ti): less than 0.06%, niobium (Nb): less than 0.06%, and boron (B): less than 0.01%.

- 4. The hot-dip galvanized or galvannealed steel sheet according to any one of claims 1 to 3, wherein said zinc coated layer is a galvannealed layer comprising zinc (Zn): 80 to 91%, iron (Fe): 8 to 15%, and aluminum (Al): not more than 1%.
- 5. The hot-dip galvanized or galvannealed steel sheet according to any one of claims 1 to 3, wherein said zinc coated layer is a galvanized layer comprising zinc: not less than 80% and aluminum: not more than 1%.
- 6. A process for producing the hot-dip galvanized or galvannealed steel sheet according to claim 4, said process comprising the steps of:

providing a cold rolled steel sheet having the chemical composition of the steel sheet substrate as defined in any one of claims 1 to 3;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, and subjecting the cooled steel sheet to hot-dip galvannealing process; and then

holding the coated steel sheet in the temperature range of 450 to 600°C for 5 sec to 2 min and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

7. A process for producing the hot-dip galvanized or galvannealed steel sheet according to claim 4, said process comprising the steps of:

providing a cold rolled steel sheet having the chemical composition of the steel sheet substrate as defined in any one of claims 1 to 3;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900° C,

then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, holding the cooled steel sheet in said cooling temperature range for not more than 10 min, and then subjecting the cooled steel sheet to hot-dip galvannealing process; and then

holding the coated steel sheet in the temperature range of 450 to 600°C for 5 sec to 2 min and then cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec.

8. A process for producing the hot-dip galvanized or galvannealed steel sheet according to claim 5, said process comprising the steps of:

providing a cold rolled steel sheet having the chemical composition of the steel sheet substrate as defined in any one of claims 1 to 3;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, and subjecting the cooled steel sheet to hot-dip galvanizing process; and then

cooling the coated steel sheet to 250°C or below at a cooling rate of not less than 5°C/sec .

9. A process for producing the hot-dip galvanized or galvannealed steel sheet according to claim 5, said process comprising the steps of:

providing a cold rolled steel sheet having the chemical composition of the steel sheet substrate as defined in any one of claims 1 to 3;

annealing the cold rolled steel sheet for 10 sec to 6 min in the intercritical temperature range of 650 to 900°C, then cooling the annealed steel sheet to 350 to 500°C at a cooling rate of 2 to 200°C/sec, holding the cooled steel sheet in said cooling temperature range for not more than 10 min, and then subjecting the cooled steel sheet to hot-dip galvanizing process; and then

cooling the coated steel sheet to 250°C or below at a

cooling rate of not less than 5°C/sec.

Fetherstonhaugh & Co. Ottawa, Canada Patent Agents